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(54) Aminoguanidines

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(56) References cited:

EP-A- 87 218 US-A- 3 317 560 US-A- 2 855 398

CHEMICAL ABSTRACTS, vol. 67, no. 13, 25
 September 1967, Columbus, Ohio, US; abstract no. 64162V, ALEMANY ET AL.: 'Potential psychotropic agents.'

 INDIAN JOURNAL OF CHEMISTRY vol. 15 B, no. 12, December 1977, NEW DELHI INDIA pages 1129 - 1132; ARYA V. P. ET AL.: 'Synthesis and CNS effects of some 2- substituted-5-acetyl-4methylpyrimidine derivatives.'

CHEMICAL ABSTRACTS, vol. 108, no. 5, 1
February 1988, Columbus, Ohio, US; abstract
no. 37353S, PITZELE ET AL.: 'potential
antisecretory antidiarrheals'

CHEMICAL ABSTRACTS, vol. 67, no. 7, 14
 August 1967, Columbus, Ohio, US; abstract no.
 32590S, EDILBERTO ET AL.: 'indol-2(or 3) ylalkyl hydrazides'

CHEMICAL ABSTRACTS, vol. 77, no. 11, 11
 September 1972, Columbus, Ohio, US; abstract no. 70181Y, OZAWA ET AL.: 'pharmacological studies of aminoquanidines.

Remarks:

The file contains technical information submitted after the application was filed and not included in this specification

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Description

The present invention relates to aminoguanidines having pharmaceutical utility, processes for their production, pharmaceutical compositions comprising them and their use as pharmaceuticals.

Aminoguanidines are disclosed in EP-A1-87218, Indian Journal of Chemistry, 15B, 1977, 1129-1132 and C.A., vol. 77, 1972, 70181y; however, these compounds comprise an aminoguanidine residue attached to a phenyl, pyrimidine or benzoyl residue.

US-A-2,855,398 describes indol-3-yl amidines having diuretic, anti-emetic and spasmolytic properties. US-A-3,317,560 discloses indol-3-yl alkylguanidines exhibiting strong spasmolytic and central depressive activities as well as dilating effects on the coronary vessels. C.A., vol. 67, 1967, 64162 v describes 1-acyl-2-(indol-3-yl-methylene) hydrazines and their in vitro activity as monoamine oxidase inhibitors. Indol-3-ylalkyl hydrazides useful in psychopharmacology against serotonin, aminooxidases, and inflammations are disclosed in C.A., vol. 67, 1967, 32590s.

It has now been found that aminoguanidines as disclosed hereafter have interesting pharmacological activity. More particularly the present invention provides a compound of formula I,

 $\begin{array}{c|c}
R_6 & Z \\
R_7 & X & Y - NH - B
\end{array}$ (I)

wherein

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R₁ is hydrogen; C₁₋₆alkyl; (C₁₋₆alkyl)carbonyl; benzoyl; or phenylC₁₋₄alkyl-carbonyl;

is hydrogen; halogen; C₁₋₆alkyl; hydroxy; nitro; amino; C₁₋₄alkylamino; C₁₋₁₀alkylcarbonylamino; C₂₋₆alkoxycarbonyl; SO₂NR_aR_b wherein each of R_a and R_b independently is hydrogen or C₁₋₆alkyl; cyano; or trimethylsilyl; C₁₋₆alkyl substituted by -SO₂-C₁₋₆alkyl, -SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂-C₁₋₆alkyl, -N(C₁₋₆alkyl)-SO₂-(C₁₋₆alkyl), -NR_aR'_b wherein R'_b is hydrogen or C₁₋₆alkyl, C₂₋₆alkoxycarbonyl or -PO(C₁₋₄alkyl)₂; carboxy; -CONR_aR_b; -PO(C₁₋₄alkyl)₂; OCONR_cR_d, wherein each of R_c and R_d independently is C₁₋₆alkyl;

35 R₆ is hydrogen or, when R₅ is OH, R₆ is hydrogen or halogen,

Z is -CR₄= wherein R₄ is hydrogen, halogen, hydroxy or C₁₋₆alkyl or, when R₅ is hydrogen or hydroxy, Z is also -N=,

R₇ is hydrogen, halogen, C₁₋₆alkyl or C₁₋₆alkoxy,

X--Y is -CR₈=N- or -CH(R₈)-NH- wherein R₈ is hydrogen or C_{1.6}alkyl, and

40 B is a radical of formula (a) or (b),

45 $A_{1} \downarrow X_{1}$ $A_{1} \downarrow X_{2}$ $A_{1} \downarrow X_{2}$ $A_{1} \downarrow X_{2}$ $A_{2} \downarrow X_{3}$ $A_{3} \downarrow X_{4}$ $A_{4} \downarrow X_{2}$ $A_{50} \downarrow X_{2}$ $A_{1} \downarrow X_{2}$ $A_{1} \downarrow X_{2}$ $A_{2} \downarrow X_{3}$ $A_{3} \downarrow X_{4}$ $A_{1} \downarrow X_{2}$ $A_{2} \downarrow X_{3}$ $A_{3} \downarrow X_{4}$ $A_{4} \downarrow X_{2}$ $A_{50} \downarrow X_{2}$ $A_{1} \downarrow X_{2}$ $A_{2} \downarrow X_{3}$ $A_{3} \downarrow X_{4}$ $A_{4} \downarrow X_{4}$ $A_{5} \downarrow X_{5}$ $A_{5} \downarrow$

wherein

wherever

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n is 1 or 2, is C=O or CH₂, is S; NR_{11} wherein R_{11} is hydrogen C_{1-6} alkylcarbonyl, benzoyl, or phenyl C_{1-4} alkyl-carbonyl; or $CR_{12}R_{13}$, wherein each of R₁₂ and R₁₃ independently is hydrogen or C_{1.4}alkyl, R₁₀ is hydrogen; C1-12alkyl; C1-6alkyl substituted by hydroxy, aryl, aryloxy, adamantyl, a heterocyclic radical, -NH₁₅-CO-R₁₆ or -NH-SO₂-aryl; C₅₋₇cycloalkyl; adamantyl; (C₁₋₁₀alkyl)carbonyl; benzoyl; phenyl(₁. 4alkyl)carbonyl; or -CONHR14, wherein R_{14} is C₁₋₁₀alkyl or C₅₋₇cycloalkyl, R₁₅ is hydrogen or C₁₋₄alkyl, and R₁₆ is C₁₋₆alkyl, C₅₋₇cycloalkyl, C₅₋₇cycloalkyl-C₁₋₄alkyl, aryl or arylC₁₋₄alkyl, winerever "aryl" appears as is or in the significances "aryloxy", ""NH-SO2-aryl" or "aryl(C1.4alkyl)" in the above defi-

nition, it is phenyl or phenyl substituted by halogen, C₁₋₄alkyl or C₁₋₆alkoxy; and "heterocyclic radical" appears in the above definition, it is pyridyl, imidazolyl, benzimidazolyl, pyrrolidinyl,

pyrrolidonyl, piperidino, pyrazinyl, perhydroindolyl or a radical of formula (c), (d) or (e)

(c) (d)

(e) 40

wherein

R₂₂ is hydrogen or C1-4alkyl,

B₁ is -CH2CH2-, -COCH2- or -(CH2)3- in which one or two H thereof can be replaced by C1-4alkyl, or 1,2-phe-

Ε is -CH2OH2-, -CH2N(R17)- or -(CH2)3- in which one or two H thereof can be replaced by C1-6alkyl, or 1,2-phenylene,

E₁ is CO or CH2,

is hydrogen or C₁₋₄alkyl, R₁₇

is CO, -CHCOOR₁₈, -CHCOR₁₉, 5,5-dimethyl-1,3-dioxan-2-ylidene or 1,3-dioxolan-2-ylidene, wherein R₁₈ is hydrogen or C₁₋₆alkyl and R₁₉ is C₁₋₆alkyl, and

n' is 0 or 1

 X_2 is $-SR_{20}$ or $-NR_3R'_{10}$ wherein R_{20} is C_{1-6} alkyl, R_3 is hydrogen or C_{1-6} alkyl and R'_{10} has one of the significances

given for R₁₀ above, or R₃ and R'₁₀ together with the nitrogen atom to which they are attached form a heterocyclic radical as defined above;

with the proviso that where B is a radical of formula (b), only one of R_{10} and R'_{10} can be other than hydrogen and X_2 can be -SR₂₀ only when R_{10} is hydrogen,

and a physiologically-hydrolysable and -acceptable ether or ester thereof when R₅ is hydroxy, in free form or in salt form.

By the term "physiologically-hydrolysable and-acceptable ethers and esters" as applied to the compounds of formula I when R_5 is hydroxy, is meant ethers in which R_5 is etherified and esters in which R_5 is esterified and which are hydrolysable under physiological conditions to yield an alcohol or acid which is physiologically acceptable, i.e. which is non-toxic at the desired dosage levels.

Examples of ether group as R_5 include e.g. C_{1-6} alkoxy; C_{1-6} alkoxy substituted by hydroxy, C_{1-4} alkoxy, acyloxy, NR_aR_b , $CONR_aR_b$ or $CSNR_aR_b$ wherein R_a , R_b and R_b are as defined above; C_{2-6} alkenyloxy.

Examples of ester groups as R_5 include e.g. acyloxy and pyridyl-carbonyloxy. When R_5 is an ester group, it is preferably pyridyl-carbonyloxy. R_5 as an ester group is preferably acyloxy or pyridyl-carbonyloxy.

In the compounds of formula I, alkyl groups and moieties may be branched or straight chain. When R_5 , R_{10} or R'_{10} are substituted alkyl, the substituent is preferably located at the end of the alkyl chain.

By halogen is preferably meant fluorine or chlorine.

When R_5 is hydroxy-substituted C_{1-6} alkoxy, it may also be alkoxy polysubstituted with hydroxy, e.g. 2,3-dihydroxy-propoxy.

Aryl is preferably phenyl or naphthyl, preferably phenyl, and may be substituted. Aryl C_{1-4} alkyl is preferably phenyl- C_{1-4} alkyl, e.g. benzyl or phenethyl, and may be substituted on the phenyl ring. Aryloxy is preferably phenoxy, and may be substituted. Aryl C_{1-6} alkoxy is e.g. benzyloxy, and may be substituted on the phenyl ring. When aryl or the aryl moiety are substituted, they may be mono-or polysubstituted, for example by halogen, C_{1-4} alkyl or C_{1-6} alkoxy. Examples are e.g. phenyl or phenyl moiety mono-or disubstituted by chlorine, methyl or methoxy.

Acyl groups or acyl moieties in acyloxy are preferably RCO, where R is C_{1-10} alkyl, C_{2-10} alkenyl, C_{5-7} cycloalkyl or aryl, preferably C_{1-10} alkyl.

When each of R_1 and R_{11} independently is C_{1-6} alkylcarbonyl, benzoyl or phenyl C_{1-4} alkylcarbonyl, it is particularly C_{1-6} alkylcarbonyl. When R_{10} is C_{1-10} alkylcarbonyl, benzoyl or phenyl C_{1-4} alkylcarbonyl, it is particularly C_{1-10} alkylcarbonyl. When R_5 is acyloxy, it is preferably R'-CO-O- where R' is C_{1-6} alkyl, phenyl or phenyl C_{1-6} alkyl.

Examples of alkyl substituted by a heterocyclic radical are e.g. 2-(2-pyrrolidone-1-yl)-ethyl, 3-benzimidazolyl-propyl. When B is a radical (b) wherein R_{10} is hydrogen and X_2 is $NR_3R'_{10}$, preferably R_3 and R'_{10} are not both hydrogen. In the compounds of formula I, the following significances are preferred either individually or in any combination or subcombination:

- 1. R₁ is H, CH₃ or C₂H₅. More preferably R₁ is H.
- 2. Z is -CR₄=.

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- R₄ is hydrogen or C₁₋₄alkyl, preferably hydrogen or methyl.
 - 4. Z is -N=, R₅ is hydroxy.
- 5. R₅ is hydrogen; hydroxy; C₁₋₆alkyl substituted by -SO₂-C₁₋₆alkyl, -SO₂NH_aR_b, -CONR_aR_b, -NH-SO₂
 C₁₋₆alkyl, -N(C₁₋₆alkyl)-SO₂-C₁₋₆alkyl or -PO(C₁₋₄alkyl)₂; acyloxy; carboxy; CONR_aR_b; -PO(C₁₋₄alkyl)₂; or OCONR_cR_d; acyloxy being C₁₋₆alkylcarbonyloxy, benzoyloxy or phenyl(C₁₋₄alkyl)carbonyloxy.
 - 6. R₇ is H or CH₃.
 - 7. X--Y is -CR₈=N-.
 - 8. R₈ is H or CH₃.
 - 9. B is a radical of formula (a), preferably a radical of formula (a) wherein X₁ is -NH-.
 - 10. B is a radical of formula (b).
 - 11. H₁₀ is hydrogen.

12. X2 is NR3R'10.

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- 13. R₃ is hydrogen or C₁₋₄alkyl.
- 14. R'_{10} is hydrogen, C_{1-10} alkyl, $(C_{1-10}$ alkyl)carbonyl, benzoyl, phenyl $(C_{1-4}$ alkyl)carbonyl, CONHR₁₄, $(CH_2)_{1.5}$ -NH-CO-R₁₆ or C_{1-6} alkyl substituted in ω by aryl, a radical of formula (d) or benzimidazolyl. More preferably R'_{10} is C_{1-12} alkyl.
- 15. R₃ and R'₁₀ together with the nitrogen atom to which they are attached are piperidino or perhydroindolyl.
- 16. The radical of formula (d) is

One group of compounds in accordance with the invention is a group of compounds of formula I wherein R_1 , H_7 , X--Y and B are as defined above, Z is -CR₄= as defined above and

is hydrogen; C₁₋₆alkyl; hydroxy, C₁₋₆alkoxy; C₁₋₆alkoxy substituted by hydroxy, C₁₋₄alkoxy, (C₁₋₆alkyl)carbonyloxy, benzoyloxy, phenyl(C₁₋₄alkyl)carbonyloxy, NH_aR'_b, CONR_aR_b or CSNR_aR_b wherein each of R_a and R_b independently is hydrogen or C₁₋₆alkyl and R'_b is hydrogen, C₁₋₆alkyl, C₃₋₇cycloalkyl, C₃₋₆alkenyl, phenyl or phenyl-C₁₋₃alkyl wherein the phenyl ring is optionally substituted; C₂₋₆alkenyloxy; pyridylcarbonyloxy; nitro; amino; C₁₋₄alkylamino; C₁₋₁₀alkylcarbonylamino; C₂₋₆alkoxycarbonyl; SO₂NR_aR_b; cyano; trimethylsilyl; C₁₋₆alkyl substituted by -SO₂-C₁₋₆alkyl, -SO₂NR_aR_b; -CONR_aR_b, -NH-SO₂-C₁₋₆alkyl, -N(C₁₋₆alkyl)-SO₂-(C₁₋₆alkyl), -NR_aR'_b, C₂₋₆alkoxycarbonyl or -PO(C₁₋₄alkyl)₂; (C₁₋₆alkyl)carbonyloxy, benzoyloxy, phenyl(C₁₋₄alkyl)carbonyloxy, carboxy; CONR_aR_b; -PO(C₁₋₄alkyl)₂; or OCONR_cR_d, wherein each of R_c and R_d independently is C₁₋₆alkyl.

Particularly preferred compounds of formula I are those wherein R_1 is H; Z is -CH= or -CCH₃=; R_7 is H or CH₃; R_5 is hydrogen, hydroxy, C_{1-6} alkoxy, C_{1-6} alkyl substituted by -SO₂- C_{1-6} alkyl, -SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂- C_{1-6} alkyl, -N(C_{1-6} alkyl)-SO₂- C_{1-6} alkyl or -PO(C_{1-4} alkyl)₂, (C_{1-6} alkyl)-carbonyloxy, benzoyloxy, phenyl(C_{1-4} alkyl)-carbonyloxy, carboxy, CONR_aR_b, PO(C_{1-4} alkyl)₂ or OCONR_cR_d.

Compounds of formula I wherein Z is -N=; R_7 is H or CH_3 ; R_5 is hydroxy or C_{1-6} alkoxy are also particularly preferred.

More particularly preferred compounds of formula I are those wherein R_1 , Z, R_7 and R_5 have one of the significances given above and B is a radical

or a radical of formula (b)

Compounds of formula I may exist in free, in salt form, in solvate or hydrate form. Salt forms may include acid addition salts and salt forms obtainable when R₅ is carboxy. Suitable pharmaceutically acceptable acid addition salt forms for use in accordance with the present invention as hereinafter described include, for example, the hydrochloride, sulfate, acetate, oxalate, maleinate and furnarate salts. When R₅ is carboxy, suitable salts are e.g. alkali metal salts such as sodium or potassium, or substituted or unsubstituted ammonium salts.

It will be appreciated that compounds of formula I, wherein X--Y is -CR₈=N- and B is a radical of formula (b')

may exist as tautomers:

wherein R_1 , R_5 , R_6 , R_8 , R_7 , Z and R'_{10} are as defined above. Compounds of formula I wherein Z is -N= and R_5 is hydroxy may also exist as tautomers:

wherein R₁, R₇, B and X---Y are as defined above.

It is to be understood that where tautomeric forms occur, the present invention embraces all tautomeric forms and their mixtures, i.e. although compounds of formula I are defined for convenience by reference to one guanidino form only or to the 5-oxo form only, the invention is not to be understood as being in any way limited by the particular nomenclature or graphic representation employed. Similar considerations apply in relation to starting materials exhibiting guanidino-tautomerism or oxy/hydroxy tautomerism as hereinafter described.

In a further aspect the present invention also provides a method for the production of compounds of formula I, which method comprises:

a) for the production of a compound of formula I wherein X--Y is -CR₈=N- reacting a compound of formula II,

$$R_{6}$$
 R_{7}
 R_{1}
 R_{8}
(III)

wherein Z, R₁, R₅, R₆, R₇ and R₈ are as defined above, with a compound of formula III,

$$H_2N-NHB$$
 (III)

wherein B is as defined above, or

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b) for the production of a compound of formula I wherein X--Y is -CHR₈-NH- hydrogenating a compound of formula I wherein Y--X is -CR₈=N-; or

c) for the production of a compound of formula I, wherein B is a radical of formula (b'), subjecting to alkylation, acylation or carbamoylation a compound of formula Ia,

$$\begin{array}{c|c}
R_5 \\
\hline
Z \\
\hline
N \\
\hline
N \\
\hline
N \\
\hline
N \\
N \\
H
\end{array}$$
(Ia)

wherein Z, R₁, R₅, R₆, R₇ and X--Y are as defined above,

d) for the production of a compound of formula I wherein R_5 is hydroxy subjecting to ether cleavage a compound of formula lb

$$\begin{array}{c|c}
R_{sa} \\
\hline
R_{1} \\
\hline
R_{1}
\end{array}$$

$$\begin{array}{c}
X \\
Y \\
\hline
N \\
\end{array}$$

$$\begin{array}{c}
Y \\
\hline
N \\
\end{array}$$

wherein

Z, R_1 , R_6 , R_7 , X-Y and B are as defined above, and R_{5a} is a cleavable ether group; or

e) for the production of a physiologically-hydrolysable and -acceptable ether or ester of a compound of formula I wherein R_5 is hydroxy etherifying or acylating a compound of formula I wherein R_5 is hydroxy and recovering compounds of formula I or a physiologically-hydrolysable and -acceptable ether or ester thereof thus obtained, in free form or in salt, solvate or hydrate form.

Process step a) may be performed analogously to known methods, e.g. conveniently in the presence of an acid, for example an inorganic acid such as hydrochloric acid or hydrobromic acid, or an organic acid such as acetic acid, p-toluene sulfonic acid or pyridinium p-toluenesulfonic acid. The reaction may conveniently be effected in the presence of a protic solvent, for example methanol, ethanol or isopropanol. The reaction may advantageously be performed at a temperature between room temperature and reflux temperature.

Process step b) may be carried out in accordance with known hydrogenation methods. When R₅ is benzyloxy it may simultaneously be cleaved to a hydroxy group.

Process step c) may be carried out by methods known in the art. Alkylation or acylation of the compounds of formula la may be conveniently effected by reaction with an alkyl, cycloalkyl or aryl halide or acyl halide or anhydride, respectively, preferably in the presence of a base, for example triethylamine or a Hunig base. Carbamoylation may be conveniently carried out, by reaction with an isocyanate such as R₁₄NCO, preferably in the presence of a solvent, for example dimethylformamide.

Process step d) may be effected analogously to methods known in the art for ether cleavage. When R_{5a} is benzyloxy, it may for example conveniently be performed by hydrogenation in the presence of a catalyst, e.g. Pd on charcoal. This reaction may be carried out in a solvent, for example an alcohol, at a temperature of from room temperature to 60° C.

R_{5a} may be alkoxy, substituted alkoxy, alkenyloxy or benzyloxy.

Process step e) may e.g. be effected by reacting a compound of formula I wherein R_5 is hydroxy with an acyl halide, preferably acyl chloride. Compounds of formula I wherein R_5 is pyridyl-carbonyloxy may be prepared by reacting a compound of formula I wherein R_5 is hydroxy with a nicotine acid halide. The reaction may conveniently be performed in a solvent such as trifluoroacetic acid or trifluoromethane sulfonic acid.

Starting materials of formula II or III are either known or may be prepared analogously to methods known and practiced in the art.

For example compounds of formula II may be prepared according to the following reaction scheme:

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R₇ NO₂ V

i. Bredereck reagent

2. Reduction

 $R_6 \stackrel{R_5}{\underset{1}{\bigvee}} Z$

R₇ H N

Modified Vilsmeier reaction

Alkylation or acylation

R₆ 7 0 R₈

H

Compounds of formula IV above may be conveniently prepared by reacting a compound of formula V with a Bredereck reagent, for example (CH₃)₂NCH(OCH₃)₂, in the absence of a solvent or in the presence of a solvent such as pyrrolidine, followed by reduction, for example with hydrogen in the presence of a palladium catalyst or with hydrazine in the presence of Raney nickel.

Compounds of formula II may conveniently be produced by submitting a compound of formula IV to a modified Vilsmeier reaction and then alkylating or acylating.

The modified Vilsmeier reaction may be performed by using a dimethyl alkylamide in the presence of POCl₃, according to methods known in the art. Alkylation or acylation may be effected in a known manner, for example in the presence of a base, e.g. K₂CO₃ or C₂H₅MgBr, in a solvent such as dimethylformamide or tetrahydrofurane.

Compounds of formula III wherein B is a radical of formula (b) wherein X_2 is other than -SR₂₀ may conveniently be prepared by reacting a compound of formula VI

wherein

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 R_{10} is as defined above and R_{21} is either -NR₃R'₁₀ or -NHNH₂

either with hydrazine when R_{21} is -NR₃R'₁₀, or with an amine of formula NHR₃R'₁₀ when R_{21} is -NHNH₂. The reaction may advantageously be carried out by heating at reflux temperature. It may be conveniently performed in a solvent, for example an alcohol such as methanol or ethanol, water or dimethylformamide, in the absence or in the presence of a basic compound, for example potassium hydroxide or carbonate.

Compounds of formula III wherein B is a radical of formula (b) wherein X₂ is -SR₂₀ may conveniently be prepared by alkylating a compound of formula VII

$$H - N + S$$

$$H - N + S$$

$$(VII)$$

with a R₂₀-yielding compound, in accordance with known methods.

Insofar as the production of the starting materials is not particularly described, the compounds are known or may be prepared analogously to methods known and practiced in the art, or as disclosed in the following examples.

The following examples are illustrative of the invention. All temperatures are in ° C.

The following abbreviations are used:

THF = tetrahydrofurane
DMF = dimethylformamide
EtOH = ethanol
MeOH = methanol

5 AcOEt = ethyl acetate
(F) = foaming
(S) = sintering

EXAMPLE 1: 5-Hydroxy-Indole-3-carboxaldehyde amino[3-(2'-pyrrolidinone-1'-yl)-propylamino]methylenehydrazone

To a solution of 0.9 g 5-hydroxy-indole-3-carboxaldehyde diaminomethylenehydrazone (4.1 mmol) in 10 ml THF containing 2 ml DMF and 0.9 ml $\rm Et_3N$ (6.2 mmol) are added at room temperature 1.3 g 3-(2'-pyrrolidinone-1'-yl)1-bromopropane (6,2 mmol). The mixture is stirred at 50 ° overnight. The mixture is then cooled to room temperature and the solvent is evaporated. The residue is chromatographied over $\rm SiO_2$ (eluant: Toluene/EtOH/NH₃ 70:30:2.5) to yield the title compound as crystals. M.p. = 158 ° (foaming).

Mass spectrum m/z (relative intensity): 343.3 (MH+, 100); 217.2 (20); 168.2 (20); 143.2 (23).

EXAMPLE 2: 5-Hydroxy-Indole-3-carboxaldehyde amino(N-methyl-N-heptylamino)methylenehydrazone

To a solution of 0.48 g 5-benzyloxy-indole-3-carboxaldehyde amino(N-methyl-N-heptylamino)methylene hydrazone .(1.1 mmol) in EtOH there is added 0.25 g 1.0 % Pd/C. The suspension is hydrogenated overnight at 45 ° C. Afterwards the suspension is filtered over silica gel, the solvent is evaporated and the residue is chromatographied over silica gel (eluant: toluene/EtOH/NH₃ 85 : 15 : 1) to yield the title compound. The pure material is crystalized from CH₂Cl₂/Hexane 2 : 8.

M.p. = 110 ° C (sintering)

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Mass spectrum m/z: 329 (M+, 40); 128 (40); 111 (60); 73 (50).

The starting materials may be produced as follows:

a) To a solution of 3.2 g 5-benzyloxy-indole-3-carboxaldehyde (12.7 mmol) and 5.0 g 1-(N-methyl-N-heptyl)-3-N-amino guanidine, hydroiodide (16.0 mmol) in 100 ml MeOH are added at 5 ° a solution of MeOH/HCl until pH = 3. After 2 hours, the solvent is evaporated and the residue taken up in AcOEt. The solution is washed with a solution of Na₂CO₂ (2N). The organic layer is dried over sodium sulfate and the solvent is evaporated. The residue is chromatographied (eluant: Toluene/EtOH/NH₃ 85:15:0.5) to yield the title compound.

Mass spectrum m/z (relative intensity): 420 (MH+, 100); 330 (7); 249 (4); 172 (16).

b) 1-(N-Methyl-N-heptyl)-3-N'-aminoguanidine, hydroiodide A solution containing 4.7 g S-methyl isothiosemicarbazide hydroiodide (20 mmol) and 3.7 ml N-methyl N-heptylamine (22 mmol) in 30 ml methanol is refluxed for 6 hours. The solution is then cooled to room temperature and the solvent is evaporated to yield 1-(N-methyl-N-heptyl)-3-N'-aminoguanidine, hydroiodide. The resulting crude material is used for the next step without further purification.

EXAMPLE 3: 5-Hydroxy-indole-3-carboxaldehyde amino(N-cyclohexylureido)methylenehydrazone

To a solution of 0.8 g 5-hydroxy-indole-3-carboxaldehyde diaminomethylenhydrazone (3.7 mmol) in 20 ml DMF is added over 5 min. at 0 $^{\circ}$ a solution of 0.5 ml cyclohexyl isocyanate (4.0 mmol) in 5 ml DMF. The solution is stirred for 4 hours. The solvent is then evaporated and the residue chromatographied (eluant: Toluene/EtOH/NH₃ 85:15:0.5) to yield the title compound as crystals. M.p. = 135 $^{\circ}$ (foaming).

Mass spectrum m/z (relative intensity): 343 (MH+, 100); 244 (50); 218 (85); 159 (33).

EXAMPLE 4: 5-Hydroxy-6-fluoro-indole-3-carboxaldehyde amino(pentylamino)methylene hydrazone

The title compound is prepared by following the procedure of Example 2. M.p. = 125 ° (foaming). 5-Benzyloxy-6-fluoro-indole-3-carboxyaldehyde used as starting material may be produced as follows:

a) 2-Nitro-4-fluoro-5-benzyloxy-toluene

To a solution of 85.6 g 2-nitro-4-fluoro-5-hydroxy-toluene (0.5 mol) in 1300 ml acetone are added at room temperature 138 g K₂CO₃ (1.0 mol). 72 ml benzyl bromide (0.6 mol) are then added dropwise over 1 hour and the resulting mixture is stirred overnight at 60 °. The solvent is evaporated and the residue taken up in AcOEt. The precipitate is removed by filtration and the solution is washed with water. The organic layer is dried over sodium sulfate, the solvent evaporated and the residue crystalized from hexane to yield 2-nitro-4-fluoro-5-benzyloxy-toluene. M.p.

= 95 °.

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Mass spectrum m/z: 261 (M+).

b) 2-[1'-(N,N-Dimethylamino)-ethan-2'-yl]-4-benzyloxy-5-fluoronitrobenzene

A solution of 126 g 2-nitro-4-fluoro-5-benzyloxy-toluene (0.48 mol) in 200 g bis-dimethylamino-t-butoxy-methane (1.15 mol) is stirred overnight at 90 $^{\circ}$. Afterwards the solvent is evaporated and the residue crystalized from MeOH to yield the b) title compound as red crystals. M.p. = 146 $^{\circ}$.

Mass spectrum m/z: 316 (M+).

c) 5-Benzyloxy-6-fluoro-Indole

A solution of 9.5 g b) compound (30.0 mmol) in 150 ml toluene and 30 ml THF containing 1 g Raney nickel is hydrogenated at room temperature. After 4 hours the suspension is filtered over hyflo and the solvent is evaporated. The residue is chromatographied under medium pressure (eluant: Toluene) to yield the b) title compound which is crystalized from hexane.

M.p. = 126° . Mass spectrum m/z: $241 (M^{+})$.

d) 5-Benzyloxy-6-fluoro-indole-3-carboxaldehyde

3.3 ml POCl $_3$ (36.0 mmol) are added dropwise at 0 ° to 14 ml DMF (180.0 mmol). After 15 min. a solution of 7.30 g of the c) compound (30 mmol) in 14 ml DMF is added dropwise over 10 min. The mixture is stirred for 1 hour at room temperature, then diluted with cold water and a solution of 7.2 g NaOH in 50 ml water is then added dropwise. The precipitate is filtered and washed with water. The resulting solid is chromatographied over SiO_2 (eluant: CH_2Cl_2) and crystalized from ether to yield the d) title compound. M.p. = 190 °.

Mass spectrum m/z (relative intensity): 269 (M+, 72); 178 (20); 150 (15); 91 (100); 65 (38).

EXAMPLE 5: 5-Hydroxy-indole-3-carboxaldehyde amino(butyrylamido)methylenehydrazone

To a solution of $0.5\,\mathrm{g}$ 5-hydroxy-indole-3-carboxaldehyde diaminomethylenehydrazone (2.3 mmol) in 5 ml DMF are added dropwise a solution of $0.4\,\mathrm{ml}$ butanoic anhydride (2.5 mmol) in 5 ml DMF. After 7 hours at room temperature the solvent is evaporated and the residue is chromatographied over SiO_2 (eluant: Toluene/EtOH/NH $_3$ 85:15:0.3). The title compound is thus obtained and precipitated from hexane. M.p. = 90 ° (foaming).

Mass spectrum m/z (relative intensity): 287 (M+, 16); 217 (8); 200 (4); 158 (30); 98 (100); 70 (46).

EXAMPLE 6: 5-Benzyloxy-indole-3-carboxaldehyde amino(pentylamino)methylenehydrazone trifluoroacetate

M.p. = 138 °.

EXAMPLE 7: 5-Hexanoyloxy-indole-3-carboxaldehyde amino(pentylamino)methylenehydrazone trifluoroacetate

To a solution of 1.0 g 5-hydroxy-indole-3-carboxaldehydeamino(pentylamino)methylenehydrazone (3.5 mmol) in 10 ml CF_3CO_2H there is added 0.72 ml hexanoylchloride (5.2 mmol) at 0 ° C. After 3 hours the reaction is quenched with 2N Na_2CO_3 and the mixture is stirred for 20 min. AcOEt is added and the organic layer is separated, washed with brine and dried over Na_2SO_4 . The solvent is evaporated and the residue is washed with either to yield the crystaline title compound.

 $M.p. = 205 ^{\circ},$

Mass spectrum m/z: 385 (M⁺, 20); 160 (30); 158 (25); 69 (100).

By following a procedure as disclosed above, the compounds of formula IA

wherein ${\rm R}_{\rm 5},\,{\rm R}_{\rm 8}$ and ${\rm R'}_{\rm 10}$ are as defined in Table I thereafter, may be prepared.

₩E.

TABLE I

5 ·	Ex.	R ₅	R ₈	R'10	M.P.
	8	OCH2OCH3	H	pentyl	108 °
10	9	OCH ₂ CH=C(CH ₃) ₂	H	pentyl	amorph
	10	ОН	Н	-(CH ₂) ₃ -NH-CO-C ₆ H ₅	179 ° (F)
15	11	OCO-N(CH ₃) ₂	Н	pentyl	90 ° (F)
	12	H	Н	pentyl	125 °
	13	осн3	H	pentyl	124 °*
20 .	14	он	H	pentyl	128 ° (F)**
	15	OH	H	Н	 247 ° hydro-
25					chloride
	. 16	он	CH ₃	Н	180 ° (F)
30	17	ОН	H	-(CH ₂) ₂ -N	165 °
		:	1	0	
35	.18	ОН	Н	CH ₃	140 ° (F)
	19	ОН	СН3	pentyl	200 °
40	20	OC2H5	Н	pentyl	114 °
	21	0-i-C ₃ H ₇	Н	pentyl	90 °
÷	22	ОН	н	3,8-dimethyl-nonyl	150 °
45	23	ОН	н	3-(p-F-phenoxy)-propyl	85 ° (F)
	24	он	Н.	-(CH ₂) ₂ -NH-CO-C ₆ P ₅	110 ° (F)
50	25	benzoyloxy	Н	pentyl	155 ° (F)
	26	-0-C0-tert.C ₄ H ₉	Н	pentyl	amorph

	Ex.	R ₅	R ₈	R'10	M.P.
5 10	27	ОН	H	O CH ₃ CH ₃ CH ₃ NH	130 ° (F)
15	28	OCH ₃	H	-(CH ₂) ₃ -NH-CO-C ₆ H ₅	amorph
	29	OCH2OCH3	H	-(CH ₂) ₃ -NH-CO-C ₆ H ₅	amorph
	30	OCH ₂ CH=C(CH ₃) ₂	н	-(CH ₂) ₃ -NH-CO-C ₆ H ₅	amorph
20	31	он	H	-S-(CH ₂) ₄ -CH ₃	190 ° ' hydro- iodide
25	32	соон .	H 	pentyl	310 ° hydro- chloride
	33	3-pyridyl-carbonyloxy	H	pentyl	95 °
30	34	он	H	3-benzamido-propyl	179 ° (F)
	35	0-C0-N(C ₂ H ₅) ₂	Н	pentyl	75 ° (F)
35	36	ОН	Н	-(CH ₂) ₃ -OH	140 ° (F)
	37	0-CH ₂ -CO-N(CH ₃) ₂	H	pentyl	160 °
	38	ОН	H	3-benzimidazol-2-yl-propyl	amorph
40	39	он	H	-(CH2)3-NH-SO2-C6H5	amorph
	40	0-CH ₂ -CH ₂ -N(CH ₃) ₂	H	pentyl	amorph
	41	0-(CH ₂) ₂ -0-CH ₃	H	pentyl	amorph
45	42	0-(CH ₂) ₂ -OH	Н	pentyl	amorph
	43	он	H	octyl	amorph
50	44	Si(CH ₃) ₃	H	pentyl	amorph
	45	isobutoxy	H	pentyl	amorph

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	Ex.	R ₅	R ₈	R'10	M.P.
5	46	OCH ₂ CS-N(CH ₃) ₂	H	pentyl	amorph
	47	он	Н	phenethyl	130 ° (F)
10	48	OH	Н	-(CH ₂) ₃ -N(CH ₃)-benzoyl	202 °
	49	2,3-di(OH)-propoxy	н	pentyl	105 ° (S)
	50	NH ₂	H:	penty]	100 ° (F)
15	51	acetoxy	H	pentyl	225 ° hydro- iodide
20	52	PO(CH ₃) ₂	H	pentyl	90 ° (F)
	53	COOCH ₃	В	pentyl	184 °
	54	CN	Н	pentyl	138 ° (F)
25	55	NO ₂	Н	pentyl	153 °
	. 56	CH ₂ -SO ₂ -NHCH ₃	Н	pentyl	98 ° (S)
30	57 ⁻	OCH ₂ OCO-t.butyl	. Н	pentyl	amorph
	58	CH ₂ -SO ₂ -NHCH ₃	Н	CO-NHC ₆ H ₁₁	180 ° (F)
	59	ОН	Н	3-phenyl-propyl	amorph
35	60	ОН	H	o-chlorophenethyl	122 ° (F)
	61	осн ₃	H	phenethyl	202 °
40	62	CH ₂ -CH ₂ -SO ₂ -CH ₃	н	pentyl	amorph
	63	CONH ₂	В	pentyl	130 ° (F)
	64	CON(CH ₃) ₂	H	pentyl	100 ° (F)

	Ex.	R ₅	R ₈	R' ₁₀	M.P.
5	65	он	Н	4-chlorophenethyl	115° (F)
	66	он	Н	3-MeO-phenethyl	120° (F)
10	67	F	H	phenethyl	212° (F)
	68	CH ₂ -N(CH ₃) ₂	H	pentyl	amorph
	69	CONH ₂	CH ₃	pentyl	246° (1)
75	70	он	H	3,4-di-Cl-phenethyl	274° (1)
	71	F	H	3-MeO-phenethyl	185° (1)
20	72	н	Н	CH ₂ CH ₂ CONH ₂	amorph (1)
	73	CH ₂ -CH ₂ -NH-SO ₂ CH ₃	Н	pentyl	105° (1;F)
	74	CH ₂ -NH-SO ₂ CH ₃	Н	pentyl	204° (1;F)
25	75	SO ₂ -NH ₂	H	pentyl	120° (F)
	76	CH(CH ₃)-OCH ₃	H	pentyl	115° (1;F)
30	77	OCH3	 H	3,4-di-Cl-phenethyl	209° (1)

* m.p. hydrogenomaleate = 190 ° C ** m.p. hydrochloride = 228 ° C (1): hydrochloride

By following a procedure as disclosed above, the compounds of formula IB

wherein $\ensuremath{\mathsf{R}}_5,\,\ensuremath{\mathsf{R}}_8$ and $\ensuremath{\mathsf{X}}$ are as defined in Table II below, may be prepared.

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TABLE II

Ex.	R ₅	R ₈	X	M.P.
78	ОН .	H.	NH.	178°(F)
79	ОН	CH ₃	NH .	240°
80	ОН	Н	CH ₂	297 ° chlorhydrate
81	ОН	н	s	165°
82	OCH₃	н	CH ₂	248 ° chlorhydrate (F)
83	CON(CH ₃) ₂	н	NH	225°
84	CH2SO2NHCH3	Ĥ"	NH	'253 °
85	CH ₂ SO ₂ NHCH ₃	СН₃	NH :	249 °
86	он .	Н	NH	140 ° (F)

20 EXAMPLE 87: 5-Hydroxy-3-[(N'-2'-imidazoline-4'-onyl)-hydrazomethyl]-indole

M.p. = 110° (F).

EXAMPLE 88:

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 $M.P. = 239^{\circ}$

By following a procedure as disclosed above, the compounds of formula IC

$$\begin{array}{c} R_{5} \\ R_{7} \\ R_{1} \\ \end{array}$$

wherein R_1 , R_3 , R_4 , R_5 , R_7 and R^\prime_{10} are as defined in Table III below, may be prepared.

TABLE III

Ex. R1 R4 R7 R5 R3 R'10 89 H H CH3 OH H pentyl 90 C2H5 H H OH H pentyl 91 H CH3 H OH H pentyl 92 H OH H H pentyl 93 H H H OH pertyl 94 H H H OH OH pentyl 95 H H H OH CH3 pentyl 96 H H H OCH3 CH3 pentyl 97 H H CH3 OCH3 CH3 pentyl 98 H H CH3 OCH3 CH3 pentyl	M.P. 130 ° (F) 144 ° 105 ° (F) 147 °
90	144 ° 105 ° (F)
91 H CH ₃ H CH H pentyl 92 H OH H H H pentyl 93 H H H OH piperidino 94 H H H OH OH pertydroindolyl 95 H H H OH CH ₃ pentyl 96 H H H OCH ₃ CH ₃ pentyl 97 H H CH ₃ OH CH ₃ pentyl	105° (F)
92 H OH H H H Pentyl 93 H H H OH piperidino 94 H H H OH Derhydroindolyl 95 H H H OH CH3 Dentyl 96 H H H OH CH3 Dentyl 97 H H CH3 OH CH3 Dentyl	
93 H H H OH piperidino 94 H H H OH perhydroindolyl 95 H H H OH CH ₃ pentyl 96 H H H OCH ₃ CH ₃ pentyl 97 H H CH ₃ OH CH ₃ pentyl	147°
94 H H H OH perhydroindolyl 95 H H H OH CH ₃ pentyl 96 H H H OCH ₃ CH ₃ pentyl 97 H H CH ₃ OH CH ₃ pentyl	
95 H H H OH CH ₃ pentyl 96 H H H OCH ₃ CH ₃ pentyl 97 H CH ₃ OH CH ₃ pentyl	164 ° (E)
96 H H H OCH ₃ CH ₃ pentyl 97 H H CH ₃ OH CH ₃ pentyl	170 ° (S)
97 H H CH ₃ OH CH ₃ pentyl	100 ° (F)
and the state of t	139°
98 H H CH ₃ OCH ₃ CH ₃ pentyl	120°(S)
	amorph
99 C ₂ H ₅ H H OH CH ₃ pentyl	138 ° (F)
100 H CH ₃ H OH H 3-benzimidazol-2-yl-prop	yl amorph
101 H H CH ₃ H H 3-benzimidazol-2-yl-prop	yl 120°(F)
102 H H OCH ₃ H H 3-benzimidazol-2-yl-prop	yl 135°(F)
103 H H CH ₃ OCR ₃ H 3,4-di-Cl-phenethyl	220 ° (1)

By following a procedure as discloseed above, the compounds of formula ID

NH NHR'10 (ID)

wherein R_7 and R'_{10} are as defined in Table IV below, may be prepared.

TABLE IV

Ex.	R ₇	R' ₁₀	M.P.
104	Н	pentyl	amorph
105	н	phenethyl	192°
106	СН₃	pentyl	195°
107	Н	CH₂CH₂NHCOC6H5	220 °
108	Н	benzyl	203°

EXAMPLE 109: (7-Azaindole)-3-carboxaldehyde amlno(pentylamino)methylenehydrazine

M.p. = 78 $^{\circ}$ (Sintering).

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EXAMPLE 110: 5-Hydroxy-6-fluoro-indole-3-carboxaldehyde amidinohydrazone

M.p. = $168 \circ (F)$.

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5-(Dimethylphosphine oxide)-indole-3-carboxaldehyde, used as starting material for the production of the compound of Example 52 may be prepared according to Example 4 d) from indol-5-dimethylphosphine oxide.

Indole-5-dimethylphosphine oxide may be prepared as follows:

EXAMPLE 111: Indoie-5-dimethylphosphine oxide

a) N-benzyl-indoline-5-(dimethylphosphine oxide)

A solution of t-BuLi in hexane (10 mmol, 1.7M) is added at - 78 ° to a solution of 5-bromo-N-benzylindoline (5 mmol) in 30 ml ether. After 10 minutes a solution of CIPO(Me)2 (10 mmol) in 10 ml THF is added thereto. The reaction is allowed to warm up to room temperature over 6 hours. Water and AcOEt are added, the organic layer is separated and the aqueous phase is extracted with AcOEt. The combined organic phases are washed with brine, dried and the solvent is evaporated. The residue is chromatographied over SiO2 (eluant : CH₂Cl₂/MeOH 95:5) to yield the a) title compound. M.p. = 180 °.

b) indoline-5-(dimethylphosphine oxide)

A solution of compound a) (1.5 mmol) in 20 ml MeOH containing 0.2 g Pd/C is hydrogenated over two hours. The solution is filtered over Hyflo and the solvent is evaporated to yield the b) title compound.

c) indole-5-(dimethylphosphine oxide)

A solution of compound b) (1.5 mmol) in 25 ml xylene containing 100 mg Pd/C is refluxed for 3 hours. The solution is filtered over Hyflo and the catalyst washed with CH₂Cl₂. The solvent is evaporated to yield the c) title compound.

 $M.p. = 195 ^{\circ}.$

The compounds of formula I and their pharmaceutically acceptable salts (hereinafter referred to as compounds of the invention) exhibit pharmaceutical activity and are, therefore, useful as pharmaceuticals.

In particular, compounds of the invention have a stimulatory effect on gastrointestinal motility as may be shown in standard test models, for example as follows:

Monopolar electrodes are implanted on the serosal side of the gut wall along the small intestine of Beagle dogs. From these electrodes, signals are fed into a preamplifier and filtered for the registration of low and high frequency potentials, in order to separate slow waves from spikes. The number of spike bursts ocurring in 2 min. periods are determined. From this the following data are extracted: duration of phase I - III, interval between 2 consecutive phase III blocks, propagation velocity. One or two cycles are recorded prior to drug administration which is done subcutaneously 10-15 min after a Phase III has passed the most distal electrodes. Control experiments are performed routinely by means of solvent administration. In fed dogs, the number of spikes per 30 min. is determined additionally. In this test the compounds of the invention stimulate myoelectric activity at dosages of the order of from about 0.001 to 10 mg/kg

Furthermore, the stimulatory effect on gastrointestinal motility of compounds of invention is also indicated e.g. by their effects on the peristaltic reflex in the isolated guinea-pig ileum.

Male guinea-pigs, 200-400g are stunned and bled. Segments of terminal ileum, 4-5 cm long, are removed and suspended as described by Trendelenburg in Arch. Exp. Path. Pharmakol., <u>81</u>, 55-129 (1917), in a 20 ml organ bath under an initial load of 1 g. The tissue is bathed with a modified Krebs solution (NaCl 118.6; CaCl₂ 2.7; KCl 4.7; KH₂PO₄ 1.2; MgSO₄ 0.1; NaHCO₃ 25.0; glucose 5.6 mM), maintained at 37°C and bubbled with 5% CO₂ in oxygen. Peristalsis is elicited for 30 s by increasing the intraluminal pressure from zero by 1 to 4 cm H₂O. Measurements are made of longitudinal muscle responses by using an isotonic force-displacement transducer and of circular muscle activity by employing a pressure transducer. The area under the curve (AUC) of peristaltic contractions is determined and concentration response curves are established by plotting the AUC representing the circular and longitudinal muscle activity. Each preparation is used as its own control, taking the peristaltic activity before the administration of the compounds to be tested as 100%. Compounds to be tested are added to the serosal side and are left in contact with the tissue for 15 min. In this test compounds of the invention have a stimulatory effect on the peristaltic activity at concentrations of the order of from about 10⁻¹⁰ M to 10⁻⁷ M.

Compounds of the invention are therefore useful for the treatment of gastrointestinal motility disorders, for example to normalize or to improve the gastric emptying and intestinal transit in subjects having a disturbed motility, e.g. gastro-

oesophageal reflux disease, decreased peristalsis of the oesophagus and/or stomach and/or small and/or large intestine, or to treat oesophagitis, gastroparesis, dyspepsia, non-ulcer dyspepsia, pseudo-obstruction, impaired colonic transit, ileus, irritable bowel syndrome, constipation, epigastric pain, postoperative gut atony, recurrent nausea and vomiting, anorexia nervosa or dyskinesias of the biliary system.

Furthermore the compounds of the invention are also indicated for use in the treatment of dyskinesias of the urinary bladder, the modulation of cortisol/aldosterone release, or for improving memory and learning.

For the above uses the required dosage will of course vary depending on the mode of administration, the particular condition to be treated and the effect desired. An indicated daily dosage is in the range of from about 0.01 to about 3 mg, e.g. from about 0.01 to about 1 mg for parenteral use, and of from about 0.1 to about 3 mg for oral use, conveniently administered once, in divided dosages 2 to 4 x/day, or in sustained release form. Unit dosage forms for oral administration accordingly comprise from about 0.0025 to about 1.5 mg active ingredient (i.e. compound or pharmaceutically acceptable salt of the invention) admixed with an appropriate solid or liquid, pharmaceutically acceptable, diluent or carrier therefor.

In accordance with the foregoing the present invention also provides:

i) A method for treating gastrointestinal motility disorders, e.g. by stimulating the motility of the gastrointestinal system, dyskinesias of the urinary bladder, modulating cortisol/aldosterone release or improving memory and learning in a subject in need thereof, which method comprises administering to said subject an effective amount of a compound of the invention or a pharmaceutically acceptable salt thereof.

It has further been found that compounds of the invention have an antiserotoninergic effect specifically at the 5-HT_4 receptors as may be shown in standard test models, for example as follows:

The isolated longitudinal muscle of the guinea pig ileum with its adhering myenteric plexus is a well established model which permits investigations of the mechanism of action of various neurotransmitters.

Method

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Male guinea pigs (200-400 g) are killed by a blow on the head and exsanguinated. A length of small intestine is removed about 2 cm from the ileo-caecal valve. The ileum is stretched over a glass rod and the mesentery is carefully removed. By strocking tangentially away from the mesenteric attachment with a wad of cotton wool, the longitudinal muscle layer is separated and stripped from the underlying circular muscle. Longitudinal muscle strips, 3-4cm length, are mounted in a 10 ml organ bath containing Tyrode solution at 37°C and bubbled with a 5% carbon dioxide in oxygen. The Tyrode solution is of the following composition (mmol/l): NaCl 137.0; CaCl₂ 1.8; KCl 2.7; MgCl₂ 1.05; NaHCO₃ 11.9; NaHPO₄ 0.4; glucose 5.6; and methysergide 0.1 µM. The strips are maintained under a resting tension of 500 mg. Contractions are recorded with an isotonic pendulum lever. After equilibration for 30 min a set concentration of carbachol is applied in 10 min intervals until a consistant reaction is achieved.

Production of the concentration/reaction curve

Non-cumulative concentration-response curves for 5-HT are established by adding increasing concentrations of the agonist to the organ bath at intervals of at least 15 min. Preceding experiments showed that the intervals were long enough to avoid tachyphylaxis. Each concentration is left in contact with the tissue for 1 min. Each strip is only used to record two concentration-response curves; the first for 5-HT alone and the second for 5-HT in the presence of a set concentration of antagonist, each strip thus serving as its own control. Antagonists are allowed to preequilibrate for at least 10 min prior to addition of 5-HT. The contractions expressed as percentage of the maximal response to 5-HT obtained from several preparations are plotted as mean values in order to obtain log-concentration-response curves. Inhibition constants are expressed in the form of pA₂ values which are graphically determined according to conventional methods (Arunlakshana et al., 1959, McKay 1978).

In this test 5-HT elicits a concentration-dependent contractile effect. 5-HT induces its major contractile effects in the longitudinal muscle strip of the guinea pig ileum by releasing substance P from nerve endings within this tissue. Its effect is mediated by two different 5-HT receptors. At low concentrations 5-HT activates a neuronal receptor which causes substance P release. The liberated substance P activates neuronal substance P receptors and this causes the release of acetylcholine which subsequently activates muscarinic receptors located on smooth muscle cells and brings about contraction. At higher concentrations 5-HT activates a second neuronal receptor which results in release of substance P to cause activation of substance P receptors on smooth muscle cells and thereby exerting contraction.

Compounds of the invention block preferentially the high affinity 5-HT₄ receptors thereby inhibiting 5-HT-induced contraction e.g. at concentrations from about 10⁻⁸ to about 10⁻⁶ mol/l. They exert less antagonistic activity at the low affinity 5-HT₃ receptor sites.

Compounds of the invention are therefore useful for the treatment of gastro-intestinal motility disorders such as tachygastria, problems of gastric emptying due to tachygastria, irritable bowel syndrome, intestinal spasms, intestinal cramps, constipation due to increased large intestinal tone, gastro-oesophageal reflex disease and dyskinesias of the biliary system.

Compounds of the invention also inhibit gastric lesions induced by necrotizing agents as indicated in standard tests, e.g. using rats with ethanol-induced gastric lesions.

The tests are carried out employing male rats (200-250 g) fasted overnight but with free access to water. The test substance is administered s.c. or orally by a metal stomach tube. Absolute ethanol is given orally 30 min after administration of the test substance and the animals are killed 1 hour later. The stomach is cut open along the greater curvature and pinned flat. Hemorrhagic erosions are quantified in two ways: area and length of the erosions.

On s.c. administration of a compound of the invention as test compound at a dosage of from ca. 0.1 µg/kg to 10 mg/kg, substantial inhibition of the gastric lesions induced by ethanol is observed compared with results for control groups receiving placebo in lieu of the test substance.

Compounds of the invention are accordingly indicated for use in the prophylactic or curative treatment of gastrointestinal disorders such as peptic ulcer diseases.

The compounds of the invention are further indicated for treating diarrhea, inflammatory diseases of the stomach and bowel, e.g. gastritis, duodenitis, including inflammatory bowel disease, nausea and vomiting. Furthermore they are also indicated for the treatment of arrhythmias, tachycardia, dyskinesia of the urinary bladder, e.g. incontinence, for reducing the occurrence of stroke, or for modulating stress responses.

For the above uses the required dosage will of course vary depending on the mode of administration, the particular condition to be treated and the effect desired. An indicated daily dosage is in the range of from about 5 μ g to about 5 mg for parenteral use, and of the order of from about 0.1 to about 100 mg for oral use, conveniently administered once, in divided dosages 2 to 4 x/day, or in sustained release form. Unit dosage forms for oral administration accordingly comprise from about 0.025 to about 50 mg of a compound of the invention admixed with an appropriate solid or liquid, pharmaceutically acceptable, diluent or carrier therefor.

In accordance with the foregoing the present invention also provides:

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ii) A method for the treatment of any of the above mentioned disorders or conditions in a subject in need thereof, which method comprises administering to said subject an effective amount of a compound of the invention or a pharmaceutically acceptable salt thereof;

Furthermore it has been found that the compounds of the invention have an antagonist effect at the central 5HT-1C receptors.

Compounds of the invention have a potent binding affinity to central 5HT-1C receptors as e.g. measured according to the method disclosed by D. Hoyer et al., Eur. J. Pharmacol., <u>118</u>, 13 - 23 (1985).

Compounds of the invention antagonise the hypolocomotion induced in rats by administration of m-chlorophenyl-piperazine (mCPP) according to the method disclosed by G.A. Kennett and G. Curzon, Br. J. Pharmacol., <u>94</u>, 137 - 147 (1988). In this test compounds of the invention counteract the mCPP induced locomotion after administration at dosages of from about 0.1 to 30 mg/kg p.o.

Compounds of the invention are therefore useful for the prophylactic treatment of migraine or for the treatment of psychiatric disorders e.g. anxiety, obsessive compulsive disorders, panic attacks, depression, bulimia, schizophrenia, situations of increased intracranial pressure and priapism.

For the above uses the required dosage will of course vary depending on the mode of administration, the particular condition to be treated and the effect desired. An indicated daily dosage is in the range of from about 0.5 to about 300 mg, conveniently administered once, in divided dosages 2 to 4 x/day, or in release form.

In accordance with the foregoing the present invention also provides:

iii) A method of prophylactic treatment of migraine or for treating psychiatric disorders in a subject in need thereof, which method comprises administering to said subject an effective amount of a compound of the invention or a pharmaceutically acceptable salt thereof.

Compounds of the invention also have an agonist effect on 5HT-1D receptors. Their binding affinity to 5HT-1D receptors has been determined e.g. according to the method disclosed by C. Waeber et al., Naunyn-Schmiedeberg's Arch. Pharmacol., 337, 595 - 601 (1988).

The agonist effect is further demonstrated in the following assay:

Anterior cerebral arteries are excised from pig brains obtained from the local slaughterhouse. Circular segments of 3-4 mm length are mounted between two L-shaped metal prongs and placed in temperature-controlled (37° C) organ baths filled with Krebs solution that is continuously gassed with 5% CO2 in oxygen. Agonist-induced vascular contrac-

tions are measured isometrically. In order to measure only 5-HT1D receptor mediated effects, ketanserin (10-7 M), which prevents contractions via 5-HT2 receptors, is added to the bath solution. Compounds of the invention induce vascular contractions at a concentration of from 10⁻¹⁰ to 10⁻⁵ M, particularly 10⁻⁹ to 10⁻⁷ M.

Compounds of the invention are therefore useful in treating conditions associated with cephalic pain, in particular in the treatment of migraine, cluster headache, chronic paroxysmal hemicrania and headache associated with vascular disorders and in alleviating the symptoms associated therewith.

For the above uses the required dosage will of course vary depending on the mode of administration, the particular condition to be treated and the effect desired. An indicated daily dosage is in the range of from about 0.5 to about 300 mg, conveniently administered once, in divided dosages 2 to 4 x/day, or in sustained release form.

In accordance with the foregoing the present invention also provides:

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iv) A method for treating conditions associated with cephalic pain, e.g. as indicated above in a subject in need thereof, which method comprises administering to said subject an effective amount of a compound of the invention or a pharmaceutically acceptable salt thereof.

The compounds of the invention may be administered by any conventional route, in particular nasally, enterally, preferably orally, e.g. in the form of tablets or capsules, or parenterally e.g. in the form of injectable solutions or suspensions or in a suppository form.

The compounds of the invention may be administered in free form or in pharmaceutically acceptable salt form. Such salts may be prepared in conventional manner and exhibit the same order of activity as the free compounds. Suitable pharmaceutically acceptable salts of the compounds of the invention include for example the hydrochlorides. Furthermore the present invention also provides:

v) A compound of the invention or a pharmaceutically acceptable salt thereof, for use as a pharmaceutical, e.g. in any of the methods as indicated above;

vi) A pharmaceutical composition comprising a compound of the invention as hereinbefore defined, or a pharmaceutically acceptable salt thereof, together with a pharmaceutically acceptable diluent or carrier therefor. Such compositions may be manufactured in conventional manner, e.g. by mixing of the ingredients.

Compounds of formula I wherein R_1 is hydrogen, R_7 is H and Z is -CH=, or wherein R_1 is H, R_7 is H, Z is -N= or -CH= and R_5 is hydroxy or C_{1-6} alkoxy have e.g. a stimulatory effect on gastrointestinal motility and are therefore useful in the method of the invention for treating motility disorders, e.g. by stimulating the motility of the gastrointestinal system as indicated above, for treating dyskinesias of the urinary bladder, modulating cortisol/aldosterone release or improving memory and learning. Compounds of Examples, 13 and 104 are preferred.

Compounds of formula I wherein R_1 and/or R_7 is other than hydrogen have e.g. an antiserotoninergic effect specifically at the 5-HT₄ receptors and inhibit gastric lesions induced by necrotizing agents and are therefore useful as an antiulcer or antimotility agent in the method of the invention for treating gastrointestinal disturbances and for the prophylactic or curative treatment of peptic ulcer diseases. They are also indicated for treating diarrhea, inflammatory diseases of the stomach and bowel, e.g. gastritis, duodenitis, including inflammatory bowel disease, nausea and vomiting, arrhythmias, tachycardia, dyskinesia of the urinary bladder, e.g. incontinence, for reducing the occurrence of stroke, or for modulating stress responses. Compounds of Examples 89, 90 and 97 are preferred.

Compounds of formula I wherein R_5 is hydrogen, hydroxy, C_{1-6} alkoxy or nitro, Z is $-CR_4$ = wherein R_4 is hydrogen, C_{1-6} alkyl, chlorine or bromine, R_7 is hydrogen or C_{1-6} alkyl, preferably those wherein B is a radical of formula (b), R'_{10} being C_{1-12} alkyl or C_{1-6} alkyl substituted by NH-CO-phenyl or benzimidazolyl, have e.g. an antagonist effect on central 5HT-1C receptors and are therefore useful in the prophylactic treatment of migraine and in the treatment of psychiatric disorders e.g. anxiety, obsessive compulsive disorders, panic attacks, depression or bulimia. Compound of Example 38 is preferred.

Compounds of formula I wherein R_5 is hydrogen, hydroxy, C_{1-6} alkoxy, carboxy, C_{2-6} alkoxycarbonyl, $CONR_aR_b$, $SO_2NH(C_{1-6}$ alkyl), C_{1-6} alkyl substituted by SO_2C_{1-6} alkyl, or $PO(C_{1-4}$ alkyl)₂, R_1 is H, H_7 is H, Z is -CH= and R_6 is hydrogen, particularly those wherein B is a radical

or a radical (b) wherein X2 is C1-12alkyl or -CONH-C6H11, have e.g. an agonist effect on 5HT-1D receptors and are

therefore useful in treating conditions associated with cephalic pain, e.g. as indicated above. Compound of Example 63 is particularly preferred.

Claims

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Claims for the following Contracting States: AT, BE, CH, DE, DK, FR, GB, IT, LI, LU, NL, PT, SE

1. A compound of formula I

 $\begin{array}{c|c}
R_{6} & Z \\
R_{7} & X & Y - NH - B
\end{array}$ (I)

wherein

 R_1 is hydrogen; C_{1-6} alkyl; $(C_{1-6}$ alkyl)carbonyl; benzoyl; or phenyl C_{1-4} alkyl-carbonyl;

is hydrogen; halogen; C_{1-6} alkyl; hydroxy; nitro; amino; C_{1-4} alkylamino; C_{1-10} alkylcarbonylamino; C_{2-6} alkoxycarbonyl; $SO_2NR_aR_b$ wherein each of R_a and R_b independently is hydrogen or C_{1-6} alkyl; cyano; or trimethylsilyl; C_{1-6} alkyl substituted by $-SO_2-C_{1-6}$ alkyl, $-SO_2NR_aR_b$, $-CONR_aR_b$, $-NH-SO_2-C_{1-6}$ alkyl, $-N(C_{1-6}$ alkyl)- $SO_2-(C_{1-6}$ alkyl), $-NR_aR_b$ wherein R_b is hydrogen or C_{1-6} alkyl, C_{2-6} alkoxycarbonyl or $-PO(C_{1-4}$ alkyl)₂; carboxy; $CONR_aR_b$; $-PO(C_{1-4}$ alkyl)₂; $COONR_cR_d$, wherein each of R_c and R_d independently is C_{1-6} alkyl;

R₆ is hydrogen or, when R₅ is OH, R₆ is hydrogen or halogen,

Z is -CR₄= wherein R₄ is hydrogen, halogen, hydroxy or C₁₋₆alkyl or, when R₅ is hydrogen or hydroxy, Z is

 R_7 is hydrogen, halogen, C_{1-6} alkyl or C_{1-6} alkoxy,

X--Y is -CR₈=N- or -CH(R₈)-NH- wherein R₈ is hydrogen or C₁₋₆alkyl, and

B is a radical of formula (a) or (b),

wherein

5 n IS 1 or 2,

X₁ is S; NR₁₁ wherein R₁₁ is hydrogen,(C₁₋₆alkyl)carbonyl, benzoyl or phenylC₁₋₄alkyl-carbonyl; or

	R ₁₀	is hydrogen; C_{1-12} alkyl; C_{1-6} alkyl substituted by hydroxy, aryl, aryloxy, adamantyl, a heterocyclic radical, -NR ₁₅ -CO-R ₁₆ or -NH-SO ₂ -aryl; C_{5-7} cycloalkyl; adamantyl; $(C_{1-10}$ alkyl)carbonyl; benzoyl; phenyl($_{1-4}$ alkyl)carbonyl; or -CONHR ₁₄ , wherein
5	R ₁₄ R ₁₅ R ₁₆ wherever	is C_{1-10} alkyl or C_{5-7} cycloalkyl, is hydrogen or C_{1-4} alkyl, and is C_{1-6} alkyl, C_{5-7} cycloalkyl, C_{5-7} cycloalkyl- C_{1-4} alkyl, aryl or aryl C_{1-4} alkyl, "aryl" appears as is or in the significances "aryloxy", "-NH- SO_2 -aryl" or "aryl(C_{1-4} alkyl)" in the above
10	wherever	definition, it is phenyl or phenyl substituted by halogen, C ₁₋₄ alkyl or C ₁₋₆ alkoxy; and "heterocyclic radical" appears in the above definition, it is pyridyl, imidazolyl, benzimidazolyl, pyrrolidinyl, pyrrolidonyl, piperidino, pyrazinyl, perhydroindolyl or a radical of formula (c), (d) or (e)
15		R ₂₂ N N-
20		B, 2
		(c) (d)
25		
30		G (CH ₂) _n ,
35		.
		(e)
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	wherein	
45	B ₁ is	hydrogen or C_{1-4} alkyl, - CH_2CH_2 -, - $COCH_2$ - or - $(CH_2)_3$ - in which one or two H thereof can be replaced by C_{1-4} alkyl, or 1,2-phelene,
	ph	-CH $_2$ CH $_2$ -, -CH $_2$ N(R $_{17}$)- or -(CH $_2$) $_3$ - in which one or two H thereof can be replaced by C $_{1-6}$ alkyl, or 1,2-enylene,
50	R ₁₇ is l G is c	CO or $\mathrm{CH_2}$, hydrogen or $\mathrm{C_{1-4}alkyl}$, CO, -CHCOR $_{19}$, 5,5-dimethyl-1,3-dioxan-2-ylidene or 1,3-dioxolan-2-ylidene, wherein $\mathrm{R_{18}}$ hydrogen or $\mathrm{C_{1-6}alkyl}$ and $\mathrm{R_{19}}$ is $\mathrm{C_{1-6}alkyl}$, and
		0 or 1
55	X ₂ is ces	$^{\circ}$ SR $_{20}$ or -NR $_3$ R' $_{10}$ wherein R $_{20}$ C $_{1-6}$ alkyl, R $_3$ is hydrogen or C $_{1-6}$ alkyl and R' $_{10}$ has one of the significans given for R $_{10}$ above, or R $_3$ and R' $_{10}$ together with the nitrogen atom to which they are attached form a terocyclic radical as defined above;

with the proviso that where B is a radical of formula (b), only one of R_{10} and R_{10} can be other than hydrogen and X_2 can be -SR₂₀ only when R_{10} is hydrogen,

and a physiologically-hydrolysable and -acceptable ether or ester thereof when R_5 is hydroxy, in free form or in salt form.

2. A compound of formula I

 R_{5} R_{7} N X Y N I

wherein

R₁, R₇, X--Y and B are as defined in claim 1,

is -CR $_4$ = wherein R $_4$ is hydrogen, halogen, hydroxy or C $_{1-6}$ alkyl, and is hydrogen; C $_{1-6}$ alkyl; hydroxy; C $_{1-6}$ alkoxy; C $_{1-6}$ alkoxy; substituted by hydroxy, C $_{1-4}$ alkoxy, (C $_{1-6}$ alkyl)carbonyloxy, benzoyloxy, phenylC $_{1-4}$ alkylcarbonyloxy, NR $_a$ R $_b$ CONR $_a$ R $_b$ or CSNR $_a$ R $_b$ wherein each of R $_a$, R $_b$ and R $_b$ independently is hydrogen or C $_{1-6}$ alkyl; C $_{2-6}$ alkenyloxy; pyridyl-carbonyloxy; nitro; amino; C $_{1-4}$ alkylamino; C $_{1-10}$ -alkylcarbonylamino; C $_{2-6}$ alkoxycarbonyl; SO $_2$ NR $_a$ R $_b$; cyano; or trimethylsilyl; C $_{1-6}$ alkyl substituted by -SO $_2$ -C $_{1-6}$ alkyl, -SO $_2$ NR $_a$ R $_b$, -CON-R $_a$ R $_b$, -NH-SO $_2$ -C $_{1-6}$ alkyl, -N(C $_{1-6}$ alkyl)-SO $_2$ -(C $_{1-6}$ alkyl), -NR $_a$ R $_b$, C $_{2-6}$ alkoxycarbonyl or -PO(C $_{1-4}$ alkyl) $_2$; (C $_{1-6}$ alkyl)carbonyloxy; benzoyloxy; phenylC $_{1-4}$ alkyl-carbonyloxy; carboxy; CONR $_a$ R $_b$; -PO(C $_{1-4}$ alkyl) $_2$; or OCONR $_a$ R $_d$, wherein each of R $_a$ and R $_d$ independently is C $_{1-6}$ alkyl,

with the proviso that where B is a radical of formula (b), only one of R_{10} and R'_{10} can be other than hydrogen and X_2 can be -SR₂₀ only when R_{10} is hydrogen, in free form or in salt form.

- 35 3. A compound according to claim 1 or 2 wherein R_1 is H, R_7 is H and Z is -CH=.
 - 4. A compound according to claim 1 wherein R_1 is H, R_7 is H, Z is -N= and R_5 is hydroxy.
- A compound according to any one of claims 1, 2 or 3 wherein R₅ is hydrogen, hydroxy, C₁₋₆alkoxy, carboxy, C₂₋₆-alkoxycarbonyl, CONR_aR_b, SO₂NH (C₁₋₆ alkyl), C₁₋₆ alkyl substituted by SO₂C₁₋₆ alkyl or PO(C₁₋₆alkyl)₂, R₁ is H, R₇ is H, Z is -CH= and R₆ is hydrogen.
 - A compound according to any one of the preceding claims wherein B is a radical of formula (b) wherein X₂ is -NR₃R'₁₀.
 - 7. 5-methoxy-indole-3-carboxaldehyde amino-(pentyl-amino)methylenehydrazone, in free form or in salt form.
- 8. A compound which is 5-hydroxy-indole-3-carboxaldehydeamino(N-cyclo-hexylureido)methylenehydrazone, 5-hydroxy-indole-3-carboxaldehyde amino(3-benzimidazol-2-yl-propylamino)methylenehydrazone, 5-carbamoyl-indole-3-carboxaldehydeamino(pentyl-amino)methylenehydrazone, 5-hydroxy-7-methyl-indole-3-carboxaldehyde amino(pentyl-amino)methylene-hydrazone, 1-ethyl-5-hydroxy-indole-3-carboxaldehyde amino(pentyl-amino)methylenehydrazone, 5-hydroxy-7-methyl-indole-3-carboxaldehyde amino (N-methyl-N-pentyl-amino)methylenehydrazone and 5-oxo-4-aza-indole-3-carboxaldehyde amino(pentyl-amino)methylenehydrazone, in free form or in salt form.
- 9. A process for the preparation of a compound of formula I as defined in claim 1, comprising
 - a) for the production of a compound of formula I wherein X-Y is -CR8=N- reacting a compound of formula II,

$$\begin{array}{c|c}
R_{6} & Z \\
R_{7} & R_{8}
\end{array}$$
(III)

wherein Z, R_1 , R_5 , R_6 , R_7 and R_8 are as defined in claim 1, with a compound of formula III,

$$H_2$$
N-NHB (III)

wherein B is as defined in claim 1; or

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- b) for the production of a compound of formula I wherein X--Y is -CHR₈-NH- hydrogenating a compound of formula I wherein Y--X is -CR₈=N-; or
- c) for the production of a compound of formula I, wherein B is a radical of formula (b'), subjecting to alkylation, acylation or carbamoylation a compound of formula Ia,

$$\begin{array}{c|c}
R_{6} & & NH \\
R_{7} & & NH \\
R_{1} & & NH_{2}
\end{array}$$
(Ia)

wherein Z, R₁, R₅, R₆, R₇ and X--Y are as defined in claim 1,

d) for the production of a compound of formula I wherein R_5 is hydroxy subjecting to ether cleavage a compound of formula Ib

$$\begin{array}{c|c}
R_{6} & Z \\
R_{7} & X & Y - NH - B
\end{array}$$
(Ib)

wherein

- Z, R₁, R₆, R₇, X--Y and B are as defined in claim 1, and R_{5a} is a cleavable ether group; or
- e) for the production of a physiologically-hydrolysable and -acceptable ether or ester of a compound of formula I wherein R_5 is hydroxy

and recovering compounds of formula I or a physiologically-hydrolysable and -acceptable ether or ester thereof

thus obtained, in free form or in salt, solvate or hydrate form.

- 10. A compound according to any one of claims 1 to 8 for use as a pharmaceutical.
- 11. A pharmaceutical composition comprising a compound according to any one of claims 1 to 8 or a pharmaceutically acceptable salt thereof, together with a pharmaceutically acceptable diluent or carrier therefor.
 - 12. Use of a compound according to any one of the claims 1 to 8 or a pharmaceutically acceptable salt thereof in the manufacture of a pharmaceutical composition for use in treating gastro-intestinal motility disorders or migraine.

Claims for the following Contracting States: ES, GR

1. A process for the production of a compound of formula I

 $\begin{array}{c|c}
R_{5} \\
\hline
R_{7} \\
\hline
R_{1}
\end{array}$ $\begin{array}{c}
X \\
\hline
\end{array}$ $\begin{array}{c}
Y \\
Y \\
Y$

wherein

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 R_1 is hydrogen; C_{1-6} alkyl; $(C_{1-6}$ alkyl)carbonyl; benzoyl; or phenyl C_{1-4} alkyl-carbonyl;

is hydrogen; halogen; C₁₋₆alkyl; hydroxy; nitro; amino; C₁₋₄alkylamino; C₁₋₁₀alkylcarbonylamino; C₂₋₆alkoxycarbonyl; SO₂NR_aR_b wherein each of R_a and R_b independently is hydrogen or C₁₋₆alkyl; cyano; or trimethylsilyl; C₁₋₆alkyl substituted by -SO₂-C₁₋₆alkyl, SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂-C₁₋₆alkyl, N(C₁₋₆alkyl)-SO₂-(C₁₋₆alkyl), -NR_aR'_b wherein R'_b is hydrogen or C₁₋₆alkyl, C₂₋₆alkoxycarbonyl or -PO(C₁₋₄alkyl)₂; carboxy; -CONR_aR_b; -PO(C₁₋₄alkyl)₂; OCONR_cR_d, wherein each of R_c and R_d independently is C₁₋₁alkyl

 R_6 is hydrogen or, when R_5 is OH, R_6 is hydrogen or halogen,

Z is -CR₄= wherein R₄ is hydrogen, halogen, hydroxy or C₁₋₆alkyl or, when R₅ is hydrogen or hydroxy, Z is also -N=

R₇ is hydrogen, halogen, C₁₋₆alkyl or C₁₋₆alkoxy,

X--Y is -CR₈=N- or -CH(R₈)-NH- wherein R₈ is hydrogen or C₁₋₆alkyl, and

B is a radical of formula (a) or (b),

$$\begin{pmatrix} (CH_2)_n \\ A_1 \\ \\ N \end{pmatrix} X_1$$

$$\begin{pmatrix} R_{10} \\ \\ N \end{pmatrix} X_2$$

$$(a)$$

$$(b)$$

wherein

	n	is 1 or 2,
	A ₁	is C=O or CH ₂ ,
	X ₁	is S; NR ₁₁ wherein R ₁₁ is hydrogen, (C ₁₋₆ alkyl)carbonyl, benzoyl, or phenylC ₁₋₄ alkyl-carbonyl; or
	•	CR ₁₂ R ₁₃ , wherein each of R ₁₂ and R ₁₃ independently is hydrogen or C ₁₋₄ alkyl,
5	R ₁₀	is hydrogen; C ₁₋₁₂ alkyl; C ₁₋₆ alkyl substituted by hydroxy, aryl, aryloxy, adamantyl, a heterocyclic radi-
		cal, -NR ₁₅ -CO-R ₁₆ or -NH-SO ₂ -aryl; C ₅₋₇ cycloalkyl; adamantyl; (C ₁₋₁₀ alkyl)carbonyl; benzoyl; phe-
		nyl(₁₋₄ alkyl)carbonyl; or -CONHR ₁₄ ,
	R ₁₄	wherein
10	R ₁₅	is C ₁₋₁₀ alkyl or C ₅₋₇ cycloalkyl, is hydrogen or C ₁₋₄ alkyl, and
	R ₁₆	is C ₁₋₆ alkyl, C ₅₋₇ cycloalkyl, C ₅₋₇ cycloalkyl-C ₁₋₄ alkyl, aryl or arylC ₁₋₄ alkyl,
	wherev	er "aryl" appears as is or in the significances "aryloxy", "-NH-SO₂-aryl" or "aryl(C₁,₄alkyl)" in the above
		definition, it is phenyl or phenyl substituted by halogen, C ₁₋₄ alkyl or C ₁₋₆ alkoxy; and
	wherev	by the same of the same of the pyridy, intiductory, benefit inductory, by the same of the
.15		inyl, pyrrolidonyl, piperidino, pyrazinyl, perhydroindolyl or a radical of formula (c), (d) or (e)
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		22
		\mathcal{E}_{1}
		1
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		(c) (d)
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		(cu ²)u,
40		(e)
45	wherein	1
		is hydrogen or C ₁₋₄ alkyl,
	B ₁	is -CH ₂ CH ₂ -, -COCH ₂ - or -(CH ₂) ₃ - in which one or two H thereof can be replaced by C ₁₋₄ alkyl, or 1,2-phenylene.
50		is $-CH_2CH_2$ -, $-CH_2N(R_{17})$ - or $-(CH_2)_3$ - in which one or two H thereof can be replaced by C_{1-6} alkyl, or 1,2-
		phenylene,
		is CO or CH ₂ ,
		is hydrogen or C ₁₋₄ alkyl,
<i>55</i>	G	is CO, -CHCOOR ₁₈ , -CHCOR ₁₉ , 5,5-dimethyl-1,3-dioxan-2-ylidene or 1,3-dioxolan-2-ylidene, wherein R ₁₈
33		is hydrogen or C ₁₋₆ alkyl and R ₁₉ is C ₁₋₆ alkyl, and is 0 or 1
		and
	X ₂	is -SR ₂₀ or -NR ₃ R' ₁₀ wherein R ₂₀ is C_{1-6} alkyl, R ₃ is hydrogen or C_{1-6} alkyl and R' ₁₀ has one of the signifi-

cances given for R_{10} above, or R_3 and R'_{10} together with the nitrogen atom to which they are attached form a heterocyclic radical as defined above;

with the proviso that where B is a radical of formula (b), only one of R_{10} and R'_{10} can be other than hydrogen and X_2 can be -SR₂₀ only when R_{10} is hydrogen.

and a physiologically-hydrolysable and -acceptable ether or ester thereof when R_5 is hydroxy, in free form or in salt form,

which process comprises

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a) for the production of a compound of formula I wherein X--Y is -CR₈=N- reacting a compound of formula II,

$$\begin{array}{c|c}
R_6 & Z & O \\
R_7 & R_1 & R_8
\end{array}$$
(II)

wherein Z, R₁, R₅, R₆, R₇ and R₈ are as defined above with a compound of formula III,

wherein B is as defined above; or

- b) for the production of a compound of formula I wherein X--Y is -CHR₈-NH- hydrogenating a compound of formula I wherein Y--X is -CR₈=N-; or
- c) for the production of a compound of formula I, wherein B is a radical of formula (b'), subjecting to alkylation, acylation or carbamoylation a compound of formula Ia,

$$R_{5}$$
 Z
 $Y-N$
 H_{1}
 (Ia)

wherein Z, R₁, R₅, R₆, R₇ and X--Y are as defined above,

d) for the production of a compound of formula I wherein R_5 is hydroxy subjecting to ether cleavage a compound of formula Ib

wherein

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Z, R_1 , R_6 , R_7 , X--Y and B are as defined above, and R_{5a} is a cleavable ether group; or

e) for the production of a physiologically-hydrolysable and -acceptable ether or ester of a compound of formula I wherein R_5 is hydroxy etherifying or acylating a compound of formula I wherein R_5 is hydroxy

and recovering compounds of formula I or a physiologically-hydrolysable and -acceptable ether or ester thereof thus obtained, in free form or in salt, solvate or hydrate form.

2. A process according to claim 1 for the production of a compound of formula I

wherein

 R_5

R₁, R₇, X--Y and B are as defined in claim 1,

is -CR₄= wherein R₄ is hydrogen, halogen, hydroxy or C₁₋₆ alkyl, and

is hydrogen; C_{1-6} alkyl; hydroxy; C_{1-6} alkoxy; C_{1-6} alkyl; C_{2-6} alkoxy; C_{1-6} alkoxy; C_{2-6} alkoxy; pyridyl-carbonyloxy; nitro; amino; C_{1-4} alkylamino; C_{1-10} -alkylcarbonylamino; C_{2-6} alkoxycarbonyl; C_{2-6} alkoxycarbonyl; C_{2-6} alkoxycarbonyl; C_{2-6} alkyl; C_{2-6} alkyl; C_{2-6} alkoxycarbonyl; C_{2-6} alkyl; C_{2-6} alkyl

with the proviso that where B is a radical of formula (b), only one of R_{10} and R'_{10} can be other than hydrogen and X_2 can be -SR₂₀ only when R_{10} is hydrogen, in free form or in salt form.

- A process according to claim 1 or 2 for the production of a compound of formula I wherein R₁ is H, R₇ is H and Z is -CH=.
- 4. A process according to claim 1 for the production of a compound of formula I wherein R₁ is H, R₇ is H, Z is -N= and R₅ is hydroxy.
 - 5. A process according to any one of claims 1, 2 or 3 for the production of a compound of formula I wherein R_5 is

hydrogen, hydroxy, C_{1-6} alkyl, carboxy, C_{2-6} alkoxycarbonyl, CONR_aR_b, SO₂NH (C₁₋₆ alkyl), C₁₋₆ alkyl substituted by SO₂C₁₋₆ alkyl or PO(C₁₋₆alkyl)₂, R₁ is H, R₇ is H, Z is -CH= and R₆ is hydrogen.

- A process according to claim 1 for the production of a compound which is 5-methoxy-indole-3-carboxaldehyde amino-(pentyl-amino)methylenehydrazone in free form or in salt form.
- 7. A process according to claim 1 for the production of a compound which is 5-hydroxy-indole-3-carboxaldehydeam-ino(N-cyclo-hexylureido)methylenehydrazone, 5-hydroxy-indole-3-carboxaldehyde amino(3-benzimidazol-2-yl-pro-pylamino)methylenehydrazone, 5-carbamoyl-indole-3-carboxaldehydeamino(pentyl-amino)methylenehydrazone, 5-hydroxy-7-methyl-indole-3-carboxaldehyde amino(pentyl-amino)methylene-hydrazone, 1-ethyl-5-hydroxy-indole-3-carboxaldehyde amino(pentyl-amino)methylenehydrazone, 5-hydroxy-7-methyl-indole-3-carboxaldehyde amino (N-methyl-N-pentyl-amino)methylenehydrazone and 5-oxo-4-aza-indole-3-carboxaldehyde amino(pentyl-amino)methylenehydrazone, in free form or in salt form.
- 15 8. Use of a compound produced according to any one of the claims 1 to 7 or a pharmaceutically acceptable salt thereof in the manufacture of a pharmaceutical composition for use in treating gastro-intestinal motility disorders or migraine.
 - 9. A compound of formula I as defined in claim 1, in free form or in salt form.
 - 10. A compound of formula I as defined in claim 2, in free form or in salt form.
 - 11. 5-methoxy-indole-3-carboxaldehyde amino-(pentyl-amino)methylenehydrazone, in free form or in salt form.
- 5 12. A pharmaceutical composition comprising a compound according to any one of claims 9 to 11 or a pharmaceutically acceptable salt thereof, together with a pharmaceutically acceptable diluent or carrier therefor.

Patentansprüche

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Patentansprüche für folgende Vertragsstaaten : AT, BE, CH, DE, DK, FR, GB, IT, LI, LU, NL, PT, SE

1. Verbindung der Formel I

$$\begin{array}{c|c}
R_{8} & Z \\
R_{7} & X & Y - NH \cdot B
\end{array}$$
(I)

worin

 R_5

R₁ steht für Wasserstoff C₁₋₆ Alkyl, (C₁₋₆Alkyl)carbonyl, Benzoyl oder Phenyl-C₁₋₄ Alkylcarbonyl,

steht für Wasserstoff Halogen, C₁₋₆ Alkyl, Hydroxy, Nitro, Amino, C₁₋₄ Alkylamino, C₁₋₁₀ Alkylcarbonyl-amino, C₂₋₆ Alkoxycarbonyl, SO₂NR_aR_b, worin jedes von R_a und R_b unabhängig für Wasserstoff oder C₁₋₆ Alkyl steht, Cyano oder Trimethylsilyl, C₁₋₆ Alkyl, das substituiert ist mit -SO₂-C₁₋₆ Alkyl, -SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂-C₁₋₆ Alkyl, -N(C₁₋₆ Alkyl)-SO₂-(C₁₋₆ Alkyl), -NR_aR'_b, worin R'_b für Wasserstoff oder C₁₋₆ Alkyl steht, C₂₋₆ Alkoxycarbonyl oder -PO(C₁₋₄ Alkyl)₂, Carboxy, -CONR_aR_b, -PO(C₁₋₄ Alkyl)₂, OCONR_cR_d, worin jedes von R_c und R_d unabhängig für C₁₋₆ Alkyl steht,

R₆ steht für Wasserstoff oder wenn R₅ für OH steht, steht R₆ für Wasserstoff oder Halogen,

Z steht f
ür -CR₄=, worin R₄ f
ür Wasserstoff, Halogen, Hydroxy oder C₁₋₆ Alkyl steht oder wenn R₅ f
ür Wasserstoff oder Hydroxy steht, steht Z auch f
ür -N=,

R₇ steht für Wasserstoff, Halogen, C₁₋₆ Alkyl oder C₁₋₆ Alkoxy.

X--Y steht für -CR8=N- oder -CH(R8)-NH-, worin R8 für Wasserstoff oder C1-6 Alkyi steht, und

B steht für einen Rest der Formel (a) oder (b),

$$\begin{pmatrix} (CH_2)_{n} \\ I \\ N \end{pmatrix} X_1$$

$$(a)$$

$$\begin{pmatrix} (CH_2)_{n} \\ I \\ N \end{pmatrix} X_2$$

$$(b)$$

worin n für 1 oder 2 steht,

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A₁ für C=O oder CH₂ steht,

X₁ steht für S, NR₁₁, worin R₁₁ für Wasserstoff, (C₁₋₆ Alkyl)carbonyl, Benzoyl oder Phenyl-C₁₋₄ Alkylcarbonyl steht, oder CR₁₂R₁₃, worin jedes von R₁₂ und R₁₃ für Wasserstoff oder C₁₋₄ Alkyl steht,

R₁₀ steht für Wasserstoff, C₁₋₁₂ Alkyl, C₁₋₆ Alkyl, das mit Hydroxy, Aryl, Aryloxy, Adamantyl, einem heterocyclischen Rest -NR₁₅-CO-R₁₆ oder -NH-SO₂-Aryl substituiert ist, C₅₋₇ Cycloalkyl, Adamantyl, (C₁₋₁₀ Alkyl)carbonyl, Benzoyl, Phenyl(C₁₋₄ Alkyl)carbonyl oder -CONHR₁₄.

worin R₁₄ für C₁₋₁₀ Alkyl oder C₅₋₇ Cycloalkyl steht,

 R_{15} für Wasserstoff oder C_{1-4} Alkyl steht und für C_{1-6} Alkyl, C_{5-7} Cycloalkyl, C_{5-7} Cycloalkyl, C_{1-4} Alkyl, Aryl oder Aryl- C_{1-4} Alkyl steht,

immer wenn "Aryl" selbst oder in den Ausdrücken "Aryloxy", -NH-SO $_2$ Aryl" oder Aryl(C_{1-4} Alkyl)" in der obigen Definition auftritt, steht es für Phenyl oder Phenyl, das mit Halogen, C_{1-4} Alkyl oder C_{1-6} Alkoxy substituiert ist, und immer wenn "heterocyclischer Rest" in der obigen Definition auftritt, steht dieser für Pyridyl, Imidazolyl, Benzimidazolyl, Pyrrolidinyl, Pyrrolidonyl, Piperidino, Pyrazinyl, Perhydroindolyl oder einen Rest der Formel (c), (d) oder (e)

(c) (d)

(e)

worin

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R₂₂ für Wasserstoff oder C₁₋₄ Alkyl steht,

B₁ für -CH₂CH₂-, -COCH₂- oder -(CH₂)₃- steht, worin ein oder zwei H hiervon durch C₁₋₄ Alkyl oder 1,2-Phe-

nylen ersetzt werden können,

E für -CH₂CH₂-, -CH₂N(R₁₇)- oder -(CH₂)₃- steht, worin eines oder zwei H hiervon durch C₁₋₆ Alkyl oder 1,2-Phenylen ersetzt werden können,

E₁ für CO oder CH₂ steht,

R₁₇ für Wasserstoff oder C₁₋₄ Alkyl steht,

G für CO, -CHCOOR₁₈, -CHCOR₁₉, 5,5-Dimethyl-1,3-dioxan-2-yliden oder 1,3-Dioxolan-2-yliden steht, worin R₁₈ für Wasserstoff oder C₁₋₆ Alkyl steht und R₁₉ für C₁₋₆ Alkyl steht und

n' für 0 oder 1 steht und

X₂ steht für -SR₂₀ oder -NH₃R'₁₀, worin R₂₀ für C₁₋₆ Alkyl steht, R₃ für Wasserstoff oder C₁₋₆ Alkyl steht und R'₁₀ eine der oben für R₁₀ angegebenen Bedeutungen hat, oder R₃ und R'₁₀ zusammen mit dem Stickstoffatom, an das sie gebunden sind, einen wie oben definierten heterocyclischen Rest bilden,

mit der Maßgabe, daß wenn B für einen Rest der Formel (b) steht, nur eines von R_{10} und R'_{10} für etwas anderes als Wasserstoff stehen kann und X_2 nur für -SR $_{20}$ stehen kann, wenn R_{10} für Wasserstoff steht, und ein physiologisch hydrolysierbarer und annehmbarer Ether oder Ester hiervon, wenn R_5 für Hydroxy steht in freier Form oder in Salzform.

50 2. Verbindung der Formel I

worin

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R₁, R₇ X--Y und B wie in Anspruch 1 definiert sind,

Tür -CR₄= steht, worin R₄ für Wasserstoff, Halogen, Hydroxy oder C₁₋₆ Alkyl steht, und steht für Wasserstoff, C₁₋₆ Alkyl, Hydroxy, C₁₋₆ Alkoxy, C₁₋₆ Alkoxy, das substituiert ist mit Hydroxy, C₁₋₄ Alkoxy, (C₁₋₆ Alkyl)carbonyloxy, Benzoyloxy, Phenyl-C₁₋₄ Alkylcarbonyloxy, NR_aR'_b, CONR_aR_b oder CSNR_aR_b, worin jedes von R_a, R_b und R'_b unabhängig für Wasserstoff oder C₁₋₆ Alkyl steht, C₂₋₆ Alkenyloxy, Pyridylcarbonyloxy, Nitro, Amino, C₁₋₄ Alkylamino, C₁₋₁₀ Alkylcarbonylamino, C₂₋₆ Alkoxycarbonyl, SO₂NR_aR_b, Cyano oder Trimethylsilyl, C₁₋₆ Alkyl, das substituiert ist mit -SO₂-C₁₋₆ Alkyl, -SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂-C₁₋₆ Alkyl, -N(C₁₋₆ Alkyl)-SO₂-(C₁₋₆ Alkyl), NR_aR'_b, C₂₋₆ Alkoxycarbonyl oder -PO(C₁₋₄ Alkyl)₂, (C₁₋₆ Alkyl)carbonyloxy, Benzoyloxy, Phenyl-C₁₋₄ Alkylcarbonyloxy, Carboxy, -CONR_aR_b, -PO(C₁₋₄ Alkyl)₂ oder OCONR_cR_d, worin jedes von R_c und R_d unabhängig für C₁₋₆ Alkyl steht.

mit der Maßgabe, daß wenn B für einen Rest der Formel (b) steht, nur eines von R_{10} und R'_{10} für etwas anderes als Wasserstoff stehen kann und X_2 nur für -SR $_{20}$ stehen kann, wenn R_{10} für Wasserstoff steht, in freier Form oder in Salzform.

- 3. Verbindung nach Anspruch 1 oder 2, worin R₁ für H steht, R₇ für H steht und Z für -CH= steht.
- 4. Verbindung nach Anspruch 1, worin R₁ für H steht, R₇ für H steht, Z für -N= steht und R₅ für Hydroxy steht.
- 5. Verbindung nach einem der Ansprüche 1, 2 oder 3, worin R₅ steht für Wasserstoff, Hydroxy, C₁₋₆ Alkoxy, Carboxy, C₂₋₆ Alkoxycarbonyl -CONR_aR_b, SO₂NH(C₁₋₆ Alkyl), C₁₋₆ Alkyl, das durch SO₂C₁₋₆ Alkyl oder -PO(C₁₋₆ Alkyl)₂ substituiert ist, R₁ für H steht, R₇ für H steht, Z für -CH= steht und R₆ für Wasserstoff steht.
- Verbindung nach einem der vorangehenden Ansprüche, worin B für einen Rest der Formel (b) steht, worin, X₂ für
 NR₃R'₁₀ steht.
 - 7. 5-Methoxyindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon in freier Form oder Salform.
- 8. Verbindung, die 5-Hydroxyindol-3-carboxaldehydamino-(N-cyclohexylureido)methylenhydrazon, 5-Hydroxyindol-3 carboxaldehydamino-(3-benzimidazol-2-yl-ppropylamino)methylenhydrazon, 5-Carbamoylindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon, 5-Hydroxy-7-methylindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon, 5-Hydroxy-7-methylindol-3-carboxaldehydamino-(N-methyl-N-pentylamino)methylenhydrazon und 5-Oxo-4-azaindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon in freier Form oder in Salzform ist.
 - 9. Verfahren zur Herstellung einer Verbindung der in Anspruch 1 definierten Formel I, gekennzeichnet durch
 - a) zur Herstellung einer Verbindung der Formel I, worin X--Y für -CR₈=N- steht, Umsetzung einer Verbindung der Formel II

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$$\begin{array}{c|c}
R_{s} & Z \\
R_{7} & R_{1}
\end{array}$$
(III)

worin Z, R₁, R₅, R₆, R₇ und R₈ wie in Anspruch 1 definiert sind, mit einer Verbindung der Formel III

worin B wie in Anspruch 1 definiert ist, oder

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- b) zur Herstellung einer Verbindung der Formel I, worin X--Y für -CHR $_8$ -NH- steht, Hydrierung einer Verbindung der Formel I, worin X--Y für -CR $_8$ -N- steht, oder
- c) zur Herstellung einer Verbindung der Formel I, worin B für einen Rest der Formel (b') steht, Durchführung einer Alkylierung, Acylierung oder Carbamoylierung mit einer Verbindung der Formel Ia,

$$\begin{array}{c|c}
R_{8} & & & NH \\
\hline
R_{7} & & & & NH \\
\hline
R_{1} & & & & &
\end{array}$$

$$\begin{array}{c|c}
X & & & & & \\
\end{array}$$

$$\begin{array}{c|c}
NH & & & & \\
NH_{2} & & & & \\
\end{array}$$

$$\begin{array}{c|c}
(Ia)
\end{array}$$

worin Z, R₁, R₅, R₆, R₇ und X--Y wie in Anspruch 1 definiert sind

d) zur Herstellung einer Verbindung der Formel I, worin R_5 für Hydroxy steht, Durchführung einer Etherspaltung mit einer Verbindung der Formel Ib

worin Z, R_1 , R_6 , R_7 , X--Y und B wie in Anspruch 1 definiert sind und R_{5a} für eine spaltbare Ethergruppe steht, oder

e) zur Herstellung eines physiologisch hydrolysierbaren und annehmbaren Ethers oder Esters einer Verbindung der Formel I, worin R_5 für Hydroxy steht, Verestern oder Acylieren einer Verbindung der Formel I, worin R_5 für Hydroxy steht,

und Gewinnen der Verbindungen der Formel I oder eines physiologisch hydrolysierbaren und annehmbaren so

erhaltenen Ethers oder Esters hiervon in freier Form oder in Form eines Salzes, Solvats oder Hydrats.

- 10. Verbindung nach einem der Ansprüche 1 bis 8 zur Verwendung als Pharmazeutikum.
- 11. Pharmazeutische Zusammensetzung, die eine Verbindung nach einem der Ansprüche 1 bis 8 oder ein pharmazeutisch annehmbares Salz hiervon zusammen mit einem pharmazeutisch annehmbaren Verdünnungsmittel oder Träger hierfür enthält.
- 12. Verwendung einer Verbindung nach einem der Ansprüche 1 bis 8 oder eines pharmazeutisch annehmbaren Salzes 10 hiervon zur Herstellung einer pharmazeutischen Zusammensetzung zur Verwendung bei der Behandlung von gastrointestinalen Motilitätsstörungen oder Migräne.

Patentansprüche für folgende Vertragsstaaten : ES. GR

1. Verfahren zur Herstellung einer Verbindung der Formel I

$$\begin{array}{c|c}
R_{5} \\
\hline
R_{7} \\
\hline
R_{1}
\end{array}$$

$$\begin{array}{c}
X \\
\end{array}$$

$$\begin{array}{c}
Y \\
\end{array}$$

$$\begin{array}{c}
X \\
\end{array}$$

$$\begin{array}{c}
Y \\
\end{array}$$

worin

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R₁ steht für Wasserstoff, C₁₋₆ Alkyl, (C₁₋₆Alkyl)carbonyl, Benzoyl oder Phenyl-C₁₋₄ Alkylcarbonyl,

 R_5 steht für Wasserstoff, Halogen, C₁₋₆ Alkyl, Hydroxy, Nitro, Amino, C₁₋₄ Alkylamino, C₁₋₁₀ Alkylcarbonylamino, C₂₋₆ Alkoxycarbonyl, SO₂NR_aR_b, worin jedes von R_a und R_b unabhängig für Wasserstoff oder C₁. 6 Alkyl steht, Cyano oder Trimethylsilyl, C1-6 Alkyl, das substituiert ist mit -SO2-C1-6 Alkyl, -SO2NRaRb, -CONR_aR_b, -NH-SO₂-C₁₋₆ Alkyl, -N(C₁₋₆ Alkyl)-SO₂-(C₁₋₆ Alkyl), -NR_aR'_b, worin R'_b für Wasserstoff oder C₁₋₆ Alkyl steht, C₂₋₆ Alkoxycarbonyl oder -PO(C₁₋₄ Alkyl)₂, Carboxy, -CONR_aR_b, -PO(C₁₋₄ Alkyl)₂, OCONR_cR_d, worin jedes von R_c und R_d unabhängig für C₁₋₆ Alkyl steht,

steht für Wasserstoff oder wenn R_5 für OH steht, steht R_6 für Wasserstoff oder Halogen, R₆

steht für -CR₄=, worin R₄ für Wasserstoff, Halogen, Hydroxy oder C₁₋₆ Alkyl steht oder wenn R₅ für Wasserstoff oder Hydroxy steht, steht Z auch für -N=,

steht für Wasserstoff, Halogen, C_{1-6} Alkyl oder C_{1-6} Alkoxy, 40 R_7

X--Y steht für -CR8=N- oder -CH(R8)-NH-, worin R8 für Wasserstoff oder C_{1-6} Alkyl steht, und

steht für einen Rest der Formel (a) oder (b),

worin n für 1 oder 2 steht,

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A₁ für C=O oder CH₂ steht,

X₁ steht für S, NR₁₁, worin R₁₁ für Wasserstoff, (C₁₋₆ Alkyl)carbonyl, Benzoyl oder Phenyl-C₁₋₄ Alkylcarbonyl steht, oder CR₁₂R₁₃, worin jedes von R₁₂ und R₁₃ unabhängig für Wasserstoff oder C₁₋₄ Alkyl steht,

R₁₀ steht für Wasserstoff, C₁₋₁₂ Alkyl, C₁₋₆ Alkyl, das mit Hydroxy, Aryl, Aryloxy, Adamantyl, einem heterocyclischen Rest -NR₁₅-CO-R₁₆ oder -NH-SO₂-Aryl substituiert ist, C₅₋₇ Cycloalkyl, Adamantyl, (C₁₋₁₀ Alkyl)carbonyl, Benzoyl, Phenyl(C₁₋₄ Alkyl)carbonyl oder -CONHR₁₄.

worin R₁₄ für C₁₋₁₀ Alkyl oder C₅₋₇ Cycloalkyl steht,

 R_{15} für Wasserstoff oder C_{1-4} . Alkyl steht und

R₁₆ für C₁₋₆ Alkyl, C₅₋₇ Cycloalkyl, C₅₋₇ Cycloalkyl-C₁₋₄ Alkyl, Aryl oder Aryl-C₁₋₄-alkyl steht,

immer wenn "Aryl" selbst oder in den Ausdrücken "Aryloxy", -NH-SO₂ Aryl" oder Aryl(C₁₋₄ Alkyl)" in der obigen Definition auftritt, steht es für Phenyl oder Phenyl, das mit Halogen, C₁₋₄ Alkyl oder C₁₋₆ Alkoxy substituiert ist, und immer wenn "heterocyclischer Rest" in der obigen Definition auftritt, steht dieser für Pyridyl, Imidazolyl, Benzimidazolyl, Pyrrolidonyl, Pyrrolidonyl, Piperidino, Pyrazinyl, Perhydroindolyl oder einen Rest der Formel (c), (d) oder (e)

E N-

(c)

(d)

(CH²)⁰,

(e)

worin

R₂₂ für Wasserstoff oder C₁₋₄ Alkyl steht,

B₁ für -CH₂CH₂-, -COCH₂- oder -(CH₂)₃- steht, worin ein oder zwei H hiervon durch C₁₋₄ Alkyl oder 1,2-Phenylen ersetzt werden können,

für -CH₂CH₂-, -CH₂N(R₁₇)- oder -(CH₂)₃- steht, worin eines oder zwei H hiervon durch C₁₋₆ Alkyl oder 1,2-Phenylen ersetzt werden k\u00f6nnen,

E₁ für CO oder CH₂ steht,

R₁₇ für Wasserstoff oder C₁₋₄ Alkyl steht,

G für CO, -CHCOOR₁₈, -CHCOR₁₉, 5,5-Dimethyl-1,3-dioxan-2-yliden oder 1,3-Dioxolan-2-yliden steht, worin R_{18} für Wasserstoff oder C_{1-6} Alkyl steht und R_{19} für C_{1-6} Alkyl steht und

n' für 0 oder 1 steht und

- X₂ steht für -SR₂₀ oder -NR₃R'₁₀, worin R₂₀ für C₁₋₆ Alkyl steht, R₃ für Wasserstoff oder C₁₋₆ Alkyl steht und R'₁₀ eine der oben für R₁₀ angegebenen Bedeutungen hat, oder R₃ und R'₁₀ zusammen mit dem Stickstoffatom, an das sie gebunden sind, einen wie oben definierten heterocyclischen Rest bilden,
- mit der Maßgabe, daß wenn B für einen Rest der Formel (b) steht, nur eines von R_{10} und R'_{10} für etwas anderes als Wasserstoff stehen kann und X_2 nur für -SR $_{20}$ stehen kann, wenn R_{10} für Wasserstoff steht, und eines physiologisch hydrolysierbaren und annehmbaren Ethers oder Esters hiervon, wenn R_5 für Hydroxy steht in freier Form oder in Salzform. wobei das Verfahren gekennzeichnet ist durch
 - a) zur Herstellung einer Verbindung der Formel I, worin X--Y für -CR₈=H- steht, Umsetzung einer Verbindung der Formel II

$$\begin{array}{c|c}
R_{0} & Z \\
R_{1} & R_{3}
\end{array}$$
(II)

worin Z, R₁, R₅, R₆, R₇ und R₈ wie oben definiert sind, mit einer Verbindung der Formel III

worin B wie oben definiert ist, oder

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- b) zur Herstellung einer Verbindung der Formel I, worin X--Y für -CHR $_8$ -NH- steht, Hydrierung einer Verbindung der Formel I, worin X--Y für -CR $_8$ =N- steht, oder
- c) zur Herstellung einer Verbindung der Formel I, worin B für einen Rest der Formel (b') steht, Durchführung einer Alkylierung, Acylierung oder Carbamoylierung mit einer Verbindung der Formel Ia,

$$\begin{array}{c|c}
R_{5} \\
\hline
R_{7} \\
\hline
R_{1}
\end{array}$$

$$\begin{array}{c}
X \\
Y \\
\hline
NH \\
H
\end{array}$$

$$\begin{array}{c}
NH \\
NH_{2}
\end{array}$$

$$\begin{array}{c}
NH \\
NH_{2}
\end{array}$$

$$\begin{array}{c}
(Ia)
\end{array}$$

- worin Z, R₁, R₅, R₆, R₇ und X--Y wie oben definiert sind
- d) zur Herstellung einer Verbindung der Formel I, worin R_5 für Hydroxy steht, Durchführung einer Etherspaltung mit einer Verbindung der Formel Ib

$$\begin{array}{c|c}
R_{6} & Z \\
\hline
R_{7} & X & Y - NH - B
\end{array}$$
(Ib)

worin Z, R₁, R₆, R₇, X--Y und B wie oben definiert sind und R_{5a} für eine spaltbare Ethergruppe steht, oder

e) zur Herstellung eines physiologisch hydrolysierbaren und annehmbaren Ethers oder Esters einer Verbindung der Formel I, worin R_5 für Hydroxy steht, Verestern oder Acylieren einer Verbindung der Formel I, worin R_5 für Hydroxy steht,

und Gewinnen der Verbindungen der Formel I oder eines physiologisch hydrolysierbaren und annehmbaren so erhaltenen Ethers oder Esters hiervon in freier Form oder in Form eines Salzes, Solvats oder Hydrats.

2. Verfahren nach Anspruch 1 zur Herstellung einer Verbindung der Formel I

worin

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R₁, R₇ X--Y und B wie in Anspruch 1 definiert sind,

Z für -CR₄= steht, worin R₄ für Wasserstoff, Halogen, Hydroxy oder C₁₋₆ Alkyl steht, und

Steht für Wasserstoff, C₁₋₆ Alkyl, Hydroxy, C₁₋₆ Alkoxy, C₁₋₆ Alkoxy, das substituiert ist mit Hydroxy, C₁₋₄ Alkoxy, (C₁₋₆ Alkyl)carbonyloxy, Benzoyloxy, Phenyl-C₁₋₄ Alkylcarbonyloxy, NR₈R'_b, CONR₈R_b oder CSNR₈R_b, worin jedes von R₈. R_b und R'_b unabhängig für Wasserstoff oder C₁₋₆ Alkyl steht, C₂₋₆ Alkenyloxy, Pyridylcarbonyloxy, Nitro, Amino, C₁₋₄ Alkylamino, C₁₋₁₀ Alkylcarbonylamino, C₂₋₆ Alkoxycarbonyl, SO₂NR₈R_b, Cyano oder Trimethylsilyl, C₁₋₆ Alkyl, das substituiert ist mit -SO₂-C₁₋₆ Alkyl, -SO₂NR₈R_b, -CONR₈R_b, -NH-SO₂-C₁₋₆ Alkyl, -N(C₁₋₆ Alkyl)-SO₂-(C₁₋₆ Alkyl), -NR₈R'_b, C₂₋₆ Alkoxycarbonyl oder -PO(C₁₋₄ Alkyl)₂, (C₁₋₆ Alkyl)carbonyloxy, Benzoyloxy, Phenyl-C₁₋₄ Alkylcarbonyloxy, Carboxy, -CONR₈R_b, -PO(C₁₋₄ Alkyl)₂ oder OCONR_cR_d, worin jedes von R_c und R_d unabhängig für C₁₋₆ Alkyl steht.

mit der Maßgabe, daß wenn B für einen Rest der Formel (b) steht, nur eines von R_{10} und R_{10} für etwas anderes als Wasserstoff stehen kann und X_2 nur für -SR $_{20}$ stehen kann, wenn R_{10} für Wasserstoff steht, in freier Form oder in Salzform.

- 55 3. Verfahren nach Anspruch 1 oder 2 zur Herstellung einer Verbindung der Formel I, worin R₁ f
 ür H steht, R₇ f
 ür H steht und Z f
 ür -CH= steht.
 - Verfahren nach Anspruch 1 zur Herstellung einer Verbindung der Formel I, worin R₁ für H steht, R₇ für H steht, Z

für -N= steht und R5 für Hydroxy steht.

- Verfahren nach einem der Ansprüche 1, 2 oder 3 zur Herstellung einer Verbindung der Formel I, worin R₅ steht für Wasserstoff, Hydroxy, C₁₋₆ Alkoxy, Carboxy, C₂₋₆ Alkoxycarbonyl -CONR_aR_b, SO₂NH(C₁₋₆ Alkyl), C₁₋₆ Alkyl, das durch SO₂C₁₋₆ Alkyl oder -PO(C₁₋₆ Alkyl)₂ substituiert ist, R₁ für H steht, R₇ für H steht, Z für -CH= steht und R₆ für Wasserstoff steht.
- 6. Verfahren nach Anspruch 1 zur Herstellung einer Verbindung, die 5-Methoxyindol-3-carboxaldehydamino(pentylamino)methylenhydrazon in freier Form oder Salzform ist.
- Verfahren nach Anspruch 1 zur Herstellung einer Verbindung, die 5-Hydroxyindol-3-carboxaldehydamino-(N-cyclohexylureido)methylenhydrazon, 5-Hydroxyindol-3-carboxaldehydamino-(3-benzimidazol-2-yl-propylamino)methylenhydrazon, 5-Carbamoylindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon, 5-Hydroxy-7-methylindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon, 1-Ethyl-5-hydroxyindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon, 5-Hydroxy-7-methylindol-3-carboxaldehydamino-(N-methyl-N-pentylamino)methylenhydrazon und 5-Oxo-4-azaindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon in freier Form oder in Salzform ist.
- 8. Verwendung einer Verbindung, die nach einem der Ansprüche 1 bis 7 hergestellt wird, oder eines pharmazeutisch annehmbaren Salzes hiervon zur Herstellung einer pharmazeutischen Zusammensetzung zur Verwendung bei der Behandlung von gastrointestinalen Motilitätsstörungen oder Migräne.
 - Verbindung der in Anspruch 1 definierten Formel I in freier Form oder Salzform.
 - Verbindung der in Anspruch 2 definierten Formel I in freier Form oder Salzform.
 - 11. 5-Methoxyindol-3-carboxaldehydamino-(pentylamino)methylenhydrazon in freier Form oder Salzform.
- 12. Pharmazeutische Zusammensetzung, die eine Verbindung nach einem der Ansprüche 9 bis 11 oder ein pharmazeutisch annehmbares Salz hiervon zusammen mit einem pharmazeutisch annehmbaren Verdünnungsmittel oder Träger hierfür enthält.

Revendications

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- Revendications pour les Etats contractants suivants : AT, BE, CH, DE, DK, FR, GB, IT, LI, LU, NL, PT, SE
 - 1. Un composé de formule I

 $\begin{array}{c|c}
R_{s} & Z \\
R_{1} & X & Y - NH - B
\end{array}$

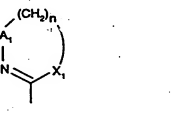
ΟÙ

R₁ signifie l'hydrogène; un groupe C₁₋₆alkyle; (C₁₋₆alkyl)carbonyle; benzoyle; ou bien phénylC₁₋₄alkyl-carbonyle;

Signifie l'hydrogène; un halogène; un groupe C₁₋₆alkyle; hydroxy; nitro; amino; C₁₋₄alkylamino; C₁₋₁₀alkylcarbonylamino; C₂₋₆alcoxycarbonyle; SO₂NR_aR_b où chacun de R_a et R_b signifie indépendamment l'hydrogène ou un groupe C₁₋₆alkyle; cyano; ou bien triméthylsilyle; un groupe C₁₋₆alkyle substitué par -SO₂- C₁₋₆alkyle, -SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂- C₁₋₆alkyle, -

 $N(C_{1-6}alkyl)-SO_2-(C_{1-6}alkyle), -NR_aR_b' où R_b' signifie l'hydrogène ou un groupe C_{1-6}alkyle, un groupe C_{2-6}alcoxycarbonyle ou -PO(C_{1-4}alkyle)_2; carboxy; -CONR_aR_b; -PO(C_{1-4}alkyle)_2; OCONR_cR_d, où chacun de R_c et R_d signifie indépendamment un groupe C_{1-6}alkyle;$

- R₆ signifie l'hydrogène ou bien, lorsque R₅ signifie OH, R₆ signifie l'hydrogène ou un halogène,
- z signifie - CR_4 = où R_4 signifie l'hydrogène, un halogène, un groupe hydroxy ou C_{1-6} alkyle ou bien, lorsque R_5 signifie l'hydrogène ou un groupe hydroxy, z signifie également -N=,
- R_7 signifie l'hydrogène, un halogène, un groupe C_{1-6} alkyle ou C_{1-6} alcoxy,
- X--Y signifie $-CR_8 = N$ ou bién $-CH(R_8)$ -NH- où R_8 signifie l'hydrogène ou bien un groupe C_{1-6} alkyle, et
- B signifie un groupe de formule (a) ou (b),



- (a) (b)
- n signifie 1 ou 2,

οù

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- A₁ signifie C=O ou bien CH₂,
- X₁ signifie S; NR₁₁ où R₁₁ signifie l'hydrogène, un groupe (C₁₋₆alkyl)carbonyle, benzoyle ou bien phényl C₁₋₄alkyl-carbonyle; ou bien CR₁₂R₁₃, où chacun de R₁₂ et R₁₃ signifie indépendamment l'hydrogène ou un groupe C₁₋₄alkyle.
- signifie l'hydrogène; un groupe C₁₋₁₂alkyle; C₁₋₆alkyle substitué par un groupe hydroxy, aryle, aryloxy, adamantyle, un groupe hétérocyclique, -NR₁₅-CO-R₁₆ ou bien -NH-SO₂-aryle; C₅₋₇cycloalkyle; adamantyle; (C₁₋₁₀alkyl)carbonyle; benzoyle; phényl(₁₋₄alkyl)carbonyle; ou bien -CONHR₁₄, où
- R₁₄ signifie un groupe C₁₋₁₀alkyle ou C₅₋₇cycloalkyle,
- R₁₅ signifie l'hydrogène ou un groupe C₁₋₄alkyle, et
- R₁₆ signifie un groupe C₁₋₆alkyle, C₅₋₇cycloalkyle, C₅₋₇cycloalkyl-C₁₋₄alkyle, aryle ou bien arylC₁₋₄alkyle,

lorsque "aryle" apparaît tel quel ou dans les significations "aryloxy", "-NH-SO₂-aryle" ou bien "aryl(C_{1-4} alkyle)" dans la définition ci-dessus, il signifie un groupe phényle ou phényle substitué par un halogène, un groupe C_{1-4} alkyle ou C_{1-6} alcoxy; et

lorsque "un groupe hérétocyclique" apparaît dans la définition ci-dessus, il signifie pyridyle, imidazolyle, benzimidazolyle, pyrrolidinyle, pyrrolidonyle, pipéridino, pyrazinyle, perhydroindolyle ou un groupe de formule (c), (d) ou (e)

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ΟÙ

R₂₂ signifie l'hydrogène ou un groupe C₁₋₄alkyle,

(e)

B₁ signifie -CH₂CH₂-, -COCH₂- ou -(CH₂)₃- dont un ou deux H peut être remplacé par un groupe C₁₋₄alkyle ou 1,2-phénylène,

signifie -CH₂CH₂-, -CH₂N(R₁₇)- ou bien -(CH₂)₃- dont un ou deux H peut être remplacé par un groupe C₁₋₆alkyle ou 1,2-phénylène,

E₁ signifie CO ou CH₂,

R₁₇ signifie l'hydrogène ou un groupe C₁₋₄alkyle,

signifie CO, -CHCOOR₁₈, -CHCOR₁₉, 5,5-diméthyl-1,3-dioxane-2-ylidène ou bien 1,3-dioxolane-2-ylidène, où R₁₈ signifie l'hydrogène ou un groupe C₁₋₆alkyle et R₁₉ signifie un groupe C₁₋₆alkyle, et

n' signifie 0 ou 1,

et

 X_2 signifie -SR₂₀ ou -NR₃R'₁₀ où R₂₀ signifie un groupe C₁₋₆alkyle, R₃ signifie l'hydrogène ou un groupe C₁₋₆alkyle et R'₁₀ a l'une des significations indiquées pour R₁₀ plus haut, ou bien R₃ et R'₁₀ forment ensemble avec l'atome d'azote auquel ils sont fixés, un groupe hérétocyclique tel que défini plus haut;

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lorsque B signifie un groupe de formule (b), seul un des symboles R_{10} et R'_{10} pouvant avoir une signification autre que l'hydrogène et X_2 pouvant signifier -SR $_{20}$ seulement lorsque R_{10} signifie l'hydrogène, et un éther ou ester physiologiquement hydrolysable et physiologiquement acceptable de ce composé lorsque R_5 signifie un groupe hydroxy, sous forme libre ou sous forme d'un sel.

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2. Un composé de formule I

οù

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R₁, R₇, X--\ Z R₅ et B sont tels que définis à la revendication 1,

signifie -CR $_4$ = où R $_4$ signifie l'hydrogène, un halogène, un groupe hydroxy ou C $_{1-6}$ alkyle, et signifie l'hydrogène; un groupe C $_{1-6}$ alkyle; hydroxy; C $_{1-6}$ alcoxy; C $_{1-6}$ alcoxy substitué par un groupe hydroxy, C $_{1-4}$ alcoxy, (C $_{1-6}$ alkyl)carbonyloxy, benzoyloxy, phénylC $_{1-4}$ alkylcarbonyloxy, NR $_4$ R $_5$, CONR $_4$ R $_5$ où chacun de R $_4$, R $_5$ et R $_5$ signifie indépendamment l'hydrogène ou un groupe C $_{1-6}$ alkyle; C $_{2-6}$ alcényloxy; pyridyl-carbonyloxy; nitro; amino; C $_{1-4}$ alkylamino; C $_{1-6}$ alkylcarbonylamino; C $_{2-6}$ alcoxycarbonyle; SO $_2$ NR $_4$ R $_5$; cyano; ou bien triméthylsilyle; C $_{1-6}$ alkyle substitué par -SO $_2$ -C $_{1-6}$ alkyle, -SO $_2$ NR $_4$ R $_5$, -CONR $_4$ R $_5$, -NH-SO $_2$ -C $_{1-6}$ alkyle, -N(C $_{1-6}$ alkyl)-SO $_2$ -(C $_{1-6}$ alkyle), -NR $_4$ R $_5$, C $_2$ -6alcoxycarbonyle ou bien -PO(C $_{1-4}$ alkyle) $_2$; (C $_{1-6}$ alkyl)-carbonyloxy; benzoyloxy; phénylC $_{1-4}$ alkyl-carbonyloxy; carboxy; CONR $_4$ R $_5$, -PO(C $_{1-4}$ alkyle) $_2$; ou bien OCONR $_5$ R $_6$, où chacun de R $_5$ et R $_6$ signifie indépendamment un groupe C $_{1-6}$ alkyle,

lorsque B signifie un groupe de formule (b), seul un des symboles R_{10} et R'_{10} pouvant avoir une signification autre que l'hydrogène et X_2 pouvant signifier -SR $_{20}$ seulement lorsque R_{10} signifie l'hydrogène, sous forme d'un sel.

- 3. Un composé selon la revendication 1 ou 2 où R₁ signifie H, R₇ signifie H et Z signifie -CH=.
- 4. Un composé selon la revendication 1 où R₁ signifie H, R₇ signifie H, Z signifie -N= et R₅ signifie un groupe hydroxy.
- 5. Un composé selon l'une quelconque des revendications 1, 2 ou 3, où R₅ signifie l'hydrogène, un groupe hydroxy, C₁₋₆alcoxy, carboxy, C₂₋₆alcoxycarbonyle, CONR_aR_b, SO₂NH (C₁₋₆alkyle), C₁₋₆alkyle substitué par SO₂C₁₋₆alkyle ou bien PO(C₁₋₆alkyle)₂, R₁ signifie H, R₇ signifie H, Z signifie -CH= et R₆ signifie l'hydrogène.
- 6. Un composé selon l'une quelconque des revendications précédentes, où B signifie un groupe de formule (b), où X₂ signifie -NR₃R₁₀.
 - La 5-méthoxy-indole-3-carboxaldéhyde amino-(pentyl-amino)méthylène-hydrazone, sous forme libre ou sous forme d'un sel.
 - 8. Un composé qui est la 5-hydroxy-indole-3-carboxaldéhydeamino-(N-cyclo-hexyluréido)méthylènehydrazone, la 5-hydroxy-indole-3-carboxaldéhyde amino(3-benzimidazole-2-yl-propylamino)méthylènehydrazone, la 5-carbamoy-lindole-3-carboxaldéhydeamino-(pentyl-amino)méthylenehydrazone, la 5-hydroxy-7-méthyl-indole-3-carboxaldéhyde amino(pentyl-amino)méthylène-hydrazone, la 1-éthyl-5-hydroxy-indole-3-carboxaldéhyde amino (N-méthyl-M-pentyl-amino)méthylènehydrazone, la 5-hydroxy-7-méthyl-indole-3-carboxaldéhyde amino (N-méthyl-M-pentyl-amino)méthylènehydrazone et la 5-oxo-4-aza-indole-3-carboxaldéhyde amino(pentyl-amino)méthylènehydrazone, sous forme libre ou sous forme d'un sel.
 - Un procédé de préparation d'un composé de formule I tel que défini à la revendication 1, selon lequel

a) pour la préparation d'un composé de formule I où X-Y signifie -CR₈=N-, on fait réagir un composé de formule II

$$\begin{array}{c|c}
R_{5} \\
\hline
R_{7} \\
\hline
R_{1}
\end{array}$$

$$\begin{array}{c}
R_{5} \\
\hline
R_{8}
\end{array}$$
(III)

où Z, R₁, R₅, R₆, R₇ et R₈ sont tels que définis à la revendication 1, avec un composé de formule III

où B est tel que défini à la revendication 1; ou bien

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b) pour la préparation d'un composé de formule I où X--Y signifie -CHR $_8$ -NH-, on hydrogène un composé de formule I où X--Y signifie -CR $_8$ -N-; ou bien

c) pour la préparation d'un composé de formule I, où B signifie un groupe de formule (b'), on introduit un groupe alkyle, acyle ou carboxy dans un composé de formule Ia,

$$\begin{array}{c|c}
R_s \\
R_7 \\
R_1
\end{array}$$

$$\begin{array}{c}
NH \\
NH_2
\end{array}$$

$$\begin{array}{c}
NH \\
NH_2
\end{array}$$

$$\begin{array}{c}
(Ia)
\end{array}$$

où Z, R_1 , R_5 , R_6 , R_7 et X--Y sont tels que définis à la revendication 1,

d) pour la préparation d'un composé de formule I où $\rm R_5$ signifie un groupe hydroxy, on soumet à une scission du groupe éther un composé de formule Ib

où Z, R₁, R₆, R₇, X--Y et B sont tels que définis à la revendication 1, et R_{5a} signifie un groupe éther scindable; ou bien

e) pour la préparation d'un éther ou d'un ester physiologiquement hydrolysable et physiologiquement acceptable d'un composé de formule I, où R_5 signifie un groupe hydroxy, on éthérifie ou on acyle un composé de formule I où R_5 signifie un groupe hydroxy,

et on récupère les composés de formule I ou un de leurs éthers ou esters physiologiquement hydrolysables et physiologiquement acceptables ainsi obtenus, sous forme libre ou sous forme d'un sel, d'un solvat ou d'un hydrate.

- 10. Un composé selon l'une quelconque des revendications 1 à 8, pour une utilisation comme médicament.
- 11. Une composition pharmaceutique comprenant un composé selon l'une quelconque des revendications 1 à 8 ou un de ses sels pharmaceutiquement acceptables, ensemble avec un diluant ou véhicule pharmaceutiquement acceptable.
- 12. L'utilisation d'un composé selon l'une quelconque des revendications 1 à 8 ou d'un de ses sels pharmaceutiquement acceptables pour la fabrication d'une composition pharmaceutique pour une utilisation dans le traitement des troubles de la motilité gastro-intestinale ou de la migraine.

Revendications pour les Etats contractants suivants : ES, GR

1. Un procédé de préparation d'un composé de formule I

$$\begin{array}{c|c}
R_s \\
\hline
R_7 \\
\hline
R_1 \\
\end{array}$$

$$\begin{array}{c}
X \\
Y \\
\hline
N \\
\end{array}$$

$$\begin{array}{c}
Y \\
\hline
N \\
\end{array}$$

ОÙ

- R₁ signifie l'hydrogène; un groupe C₁₋₆alkyle; (C₁₋₆alkyl)carbonyle; benzoyle; ou bien phénylC₁₋₄alkyl-carbonyle;
- signifie l'hydrogène; un halogène; un groupe C_{1-6} alkyle; hydroxy; nitro; amino; C_{1-4} alkylamino; C_{1-10} alkylcarbonylamino; C_{2-6} alcoxycarbonyle; $SO_2NR_aR_b$ où chacun de R_a et R_b signifie indépendamment l'hydrogène ou un groupe C_{1-6} alkyle; cyano; ou bien triméthylsilyle; un groupe C_{1-6} alkyle substitué par SO_2 C_{1-6} alkyle, $-SO_2NR_aR_b$, $-CONR_aR_b$, $-NH-SO_2$ C_{1-6} alkyle, $-N(C_{1-6}$ alkyl)- SO_2 -(C_{1-6} alkyle), $-NR_aR_b$ où R_b signifie l'hydrogène ou un groupe C_{1-6} alkyle, un groupe C_{2-6} alcoxycarbonyle ou $-PO(C_{1-4}$ alkyle) $_2$; carboxy; $-CONR_aR_b$; $-PO(C_{1-4}$ alkyle) $_2$; $OCONR_cR_d$, où chacun de R_c et R_d signifie indépendamment un groupe C_{1-6} alkyle;
- R₆ signifie l'hydrogène ou bien, lorsque R₅ signifie OH, R₆ signifie l'hydrogène ou un halogène,
- Z signifie -CR₄ = où R₄ signifie l'hydrogène, un halogène, un groupe hydroxy ou C₁₋₆alkyle ou bien, lorsque R₅ signifie l'hydrogène ou un groupe hydroxy, Z signifie également -N=,
- R₇ signifie l'hydrogène, un halogène, un groupe C₁₋₆alkyle ou C₁₋₆alcoxy,
- X--Y signifie - CR_8 = N- ou bien - $CH(R_8)$ -NH- où R_8 signifie l'hydrogène ou bien un groupe C_{1-6} alkyle, et
- B signifie un groupe de formule (a) ou (b),

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οù

n signifie 1 ou 2,

A₁ signifie C=O ou bien CH₂,

X₁ signifie S; NR₁₁ où R₁₁ signifie l'hydrogène, un groupe (C₁₋₆alkyl)carbonyle, benzoyle ou bien phényl C₁₋₄alkyl-carbonyle; ou bien CR₁₂R₁₃, où chacun de R₁₂ et R₁₃ signifie indépendamment l'hydrogène ou un groupe C₁₋₄alkyle,

R₁₀ signifie l'hydrogène; un groupe C₁₋₁₂alkyle; C₁₋₆alkyle substitué par un groupe hydroxy, aryle, aryloxy, adamantyle, un groupe hétérocyclique, -NR₁₅-CO-R₁₆ ou bien -NH-SO₂-aryle; C₅₋₇cycloalkyle; adamantyle; (C₁₋₁₀alkyl)carbonyle; benzoyle; phényl(₁₋₄alkyl)carbonyle; ou bien -CONHR₁₄, où

R₁₄ signifie un groupe C₁₋₁₀alkyle ou C₅₋₇cycloalkyle,

R₁₅ signifie l'hydrogène ou un groupe C₁₋₄alkyle, et

R₁₆ signifie un groupe C₁₋₆alkyle, C₅₋₇cycloalkyle, C₅₋₇cycloalkyl-C₁₋₄alkyle, aryle ou bien arylC₁₋₄alkyle,

lorsque "aryle" apparaît tel quel ou dans les significations "aryloxy", "-NH-SO₂-aryle" ou bien "aryl $(C_{1.4}$ alkyle)" dans la définition ci-dessus, il signifie un groupe phényle ou phényle substitué par un halogène, un groupe $C_{1.4}$ alkyle ou $C_{1.6}$ alcoxy; et

lorsque "un groupe hérétocyclique" apparaît dans la définition ci-dessus, il signifie pyridyle, imidazolyle, benzimidazolyle, pyrrolidinyle, pyrrolidonyle, pipéridino, pyrazinyle, perhydroindolyle ou un groupe de formule (c), (d) ou (e)

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$$R_{22} \stackrel{\text{O}}{\longrightarrow} N$$

$$E \stackrel{\text{N}}{\longrightarrow} N$$

$$(d)$$

G (CH2)

(e)

•

R₂₂ signifie l'hydrogène ou un groupe C₁₋₄alkyle,
 B₁ signifie -CH₂CH₂-, -COCH₂- ou -(CH₂)₃- dont un ou deux H peut être remplacé par un groupe C₁₋₄alkyle ou 1.2-phénylàne

signifie -CH₂CH₂-, -CH₂N(R₁₇)- ou bien -(CH₂)₃- dont un ou deux H peut être remplacé par un groupe C₁.

6alkyle ou 1,2-phénylène,

E₁ signifie CO ou CH₂,

R₁₇ signifie l'hydrogène ou un groupe C₁₋₄alkyle,

G signifie CO, -CHCOOR₁₈, -CHCOR₁₉, 5,5-diméthyl-1,3-dioxane-2-ylidène ou bien 1,3-dioxolane-2-ylidène, où R₁₈ signifie l'hydrogène ou un groupe C₁₋₆alkyle et R₁₉ signifie un groupe C₁₋₆alkyle, et

n' signifie 0 ou 1,

et

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οù

X₂ signifie -SR₂₀ ou -NR₃R'₁₀ où R₂₀ signifie un groupe C₁₋₆alkyle, R₃ signifie l'hydrogène ou un groupe C₁₋₆alkyle et R'₁₀ a l'une des significations indiquées pour R₁₀ plus haut, ou bien R₃ et R'₁₀ forment ensemble avec l'atome d'azote auquel ils sont fixés, un groupe hérétocyclique tel que défini plus haut;

lorsque B signifie un groupe de formule (b), seul un des symboles R_{10} et R'_{10} pouvant avoir une signification autre que l'hydrogène et X_2 pouvant signifier -S R_{20} seulement lorsque R_{10} signifie l'hdrogène, et un éther ou ester physiologiquement hydrolysable et physiologiquement acceptable de ce composé lorsque R_5 signifie un groupe hydroxy, sous forme libre ou sous forme d'un sel, procédé selon lequel

a) pour la préparation d'un composé de formule I où X--Y signifie -CR₈=N-, on fait réagir un composé de formule II

$$\begin{array}{c|c}
R_s \\
R_7 \\
R_1
\end{array}$$

$$\begin{array}{c}
R_1 \\
R_2
\end{array}$$

$$\begin{array}{c}
R_1 \\
R_2
\end{array}$$

$$\begin{array}{c}
R_2 \\
R_3
\end{array}$$

$$\begin{array}{c}
R_3 \\
R_4
\end{array}$$

où Z, R₁, R₅, R₆, R₇ et R₈ sont tels que définis plus haut, avec un composé de formule III

où B est tel que défini plus haut; ou bien

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b) pour la préparation d'un composé de formule I où X--Y signifie -CHR $_8$ -NH-, on hydrogène un composé de formule I où X--Y signifie -CR $_8$ -N-; ou bien

c) pour la préparation d'un composé de formule I, où B signifie un groupe de formule (b'), on introduit un groupe alkyle, acyle ou carboxy dans un composé de formule la,

$$\begin{array}{c|c}
R_{8} & Z & NH \\
R_{7} & X & Y - NH_{2}
\end{array}$$
(Ia)

où Z, R₁, R₅, R₆, R₇ et X--Y sont tels que définis plus haut,

d) pour la préparation d'un composé de formule I où R_5 signifie un groupe hydroxy, on soumet à une scission du groupe éther un composé de formule Ib

$$R_{sa}$$

$$Z$$

$$R_{t}$$

$$X - NH - B$$

$$R_{t}$$

$$(1b)$$

Z, R_1 , R_6 , R_7 , X--Y et B sont tels que définis plus haut, et R_{5a} signifie un groupe éther scindable; ou bien

e) pour la préparation d'un éther ou d'un ester physiologiquement hydrolysable et physiologiquement acceptable d'un composé de formule I, où R_5 signifie un groupe hydroxy, on éthérifie ou on acyle un composé de formule I où R_5 signifie un groupe hydroxy,

et on récupère les composés de formule I ou un de leurs éthers ou esters physiologiquement hydrolysables et phy-

siologiquement acceptables ainsi obtenus, sous forme libre ou sous forme d'un sel, d'un solvat ou d'un hydrate.

2. Un procédé selon la revendication 1 pour la préparation d'un composé de formule I

οù

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R₁, R₇, X--Y Z et B sont tels que définis à la revendication 1,

signifie -CR₄= où R₄ signifie l'hydrogène, un halogène, un groupe hydroxy ou C_{1-6} alkyle, et signifie l'hydrogène; un groupe C_{1-6} alkyle; hydroxy; C_{1-6} alcoxy; C_{1-6} alcoxy substitué par un groupe hydroxy, C_{1-4} alcoxy, $(C_{1-6}$ alkyl)carbonyloxy, benzoyloxy, phényl C_{1-4} alkylcarbonyloxy, NR_aR'_b, CONR_aR_b ou CSNR_aR_b où chacun de R_a, R_b et R'_b signifie indépendamment l'hydrogène ou un groupe C_{1-6} alkyle; C_{2-6} alcényloxy; pyridyl-carbonyloxy; nitro; amino; C_{1-4} alkylamino; C_{1-10} -alkylcarbonylamino; C_{2-6} alcoxycarbonyle; SO₂NR_aR_b; cyano; ou bien triméthylsilyle; C_{1-6} alkyle substitué par -SO₂- C_{1-6} alkyle, -SO₂NR_aR_b, -CONR_aR_b, -NH-SO₂- C_{1-6} alkyle, -N(C_{1-6} alkyl)-SO₂-(C_{1-6} alkyle), -NR_aR'_b, C_{2-6} alcoxycarbonyle ou bien -PO(C_{1-4} alkyle)₂; (C_{1-6} alkyl)carbonyloxy; benzoyloxy; phényl C_{1-4} alkyl-carbonyloxy; carboxy; CONR_aR_b, -PO(C_{1-4} alkyle)₂; ou bien OCONR_cR_d, où chacun de R_c et R_d signifie indépendamment un groupe C_{1-6} alkyle,

lorsque B signifie un groupe de formule (b), seul un des symboles R_{10} et R'_{10} pouvant avoir une signification autre que l'hydrogène et X_2 pouvant signifier -S R_{20} seulement lorsque R_{10} signifie l'hydrogène, sous forme libre ou sous forme d'un sel.

- Un procédé selon la revendication 1 ou 2 pour la préparation d'un composé de formule I où R₁ signifie H, R₇ signifie
 H et Z signifie -CH=.
 - 4. Un procédé selon la revendication 1 pour la préparation d'un composé de formule I où R₁ signifie H, R₇ signifie H, Z signifie -N= et R₅ signifie un groupe hydroxy.
- 5. Un procédé selon l'une quelconque des revendications 1, 2 ou 3, pour la préparation d'un composé de formule I où R₅ signifie l'hydrogène, un groupe hydroxy, C₁₋₆alcoxy, carboxy, C₂₋₆-alcoxycarbonyle, CONR_aR_b, SO₂NH (C₁₋₆alkyle), C₁₋₆alkyle substitué par SO₂C₁₋₆alkyle ou bien PO(C₁₋₆alkyle)₂, R₁ signifie H, R₇ signifie H, Z signifie CH= et R₆ signifie l'hydrogène.
- 45 6. Un procédé selon la revendication 1 pour la préparation d'un composé qui est la 5-méthoxy-indole-3-carboxaldéhyde amino-(pentyl-amino)méthylènehydrazone, sous forme libre ou sous forme d'un sel.
 - 7. Un procédé selon la revendication 1 pour la préparation d'un composé qui est la 5-hydroxy-indole-3-carboxaldéhydeamino-(N-cyclo-hexyluréido)méthylènehydrazone, la 5-hydroxy-indole-3-carboxaldéhyde amino(3-benzimidazole-2-yl-propylamino)méthylènehydrazone, la 5-carbamoyl-indole-3-carboxaldéhydeamino(pentyl-amino)méthylènehydrazone, la 5-hydroxy-7-méthyl-indole-3-carboxaldéhyde amino-(pentyl-amino)méthylènehydrazone, la 5-hydroxy-7-méthyl-indole-3-carboxaldéhyde amino(pentyl-amino)méthylènehydrazone et la 5-oxo-4-aza-indole-3-carboxaldéhyde amino(pentyl-amino)méthylènehydrazone, sous forme libre ou sous forme d'un sel.
 - 8. L'utilisation d'un composé préparé selon l'une quelconque des revendications 1 à 7 ou d'un de ses sels pharmaceutiquement acceptables, pour la fabrication d'une composition pharmaceutique pour une utilisation dans le traitement des troubles de la motilité gastro-intestinale ou de la migraine.

- 9. Un composé de formule I tel que défini à la revendication 1, sous forme libre ou sous forme d'un sel.
- 10. Un composé de formule I tel que défini à la revendication 2, sous forme libre ou sous forme d'un sel.

- La 5-méthoxy-indole-3-carboxaldéhyde amino-(pentyl-amino)méthylènehydrazone, sous forme libre ou sous forme d'un sel.
 - 12. Une composition pharmaceutique comprenant un composé selon l'une quelconque des revendications 9 à 11 ou un de ses sels pharmaceutiquement acceptables, ensemble avec un diluant ou véhicule pharmaceutiquement acceptable.